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(54) Title: GENE EXPRESSION PROFILES IN NORMAL AND CANCER CELLS

(57) Abstract

As a step towards understanding the complex differences between normal and cancer cells, gene expression patterns were examined in gastrointestinal tumors. More than 300,000 transcripts derived from at least 45,000 different genes were analyzed. Although extensive similarity was noted between the expression profiles, more than 500 transcripts that were expressed at significantly different levels in normal and neoplastic cells were identified. These data provide insights into the extent of expression differences underlying malignancy and reveal genes that are useful as diagnostic or prognostic markers.

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Gene Expression Profiles in Normal and Cancer Cells

This invention was made with support from the National Institutes of Health, Grant No. GM07309, CA57345, and CA62924. The U.S. government therefore retains certain rights in the invention.

TECHNICAL FIELD OF THE INVENTION

This invention is related to the diagnosis of cancer, and tools for carrying out such diagnosis.

BACKGROUND OF THE INVENTION

Much of cancer research over the past 50 years has been devoted to the analyses of genes that are expressed differently in tumor cells compared to their normal counterparts. Although hundreds of studies have pointed out differences in the expression of one or a few genes, no comprehensive study of gene expression in the cancer cell has been reported. It is therefore not known how many genes are expressed differentially in tumor versus normal cells, whether the bulk of these differences are cell autonomous rather than being dependent on the tumor microenvironment, and whether most differences are cell-type specific or tumor specific. Thus there is a need in the art for information on the molecular changes that occur in cells during cancer development and progression.

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SUMMARY OF THE INVENTION

According to one embodiment of the invention, a method is provided for diagnosing colon cancer in a sample suspected of being neoplastic. The method comprises the steps of:

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comparing the level of at least one transcript in a first sample of a tissue to a second sample, wherein the first sample is of a colonic tissue suspected of being neoplastic and the second sample is of a normal human colonic tissue, and wherein the transcript is identified by a tag selected from the group consisting of those shown in Table 3;

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identifying the first sample as neoplastic when the level of the at least one transcript is found to be lower in the first sample than in the second sample.

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According to another embodiment of the invention, another method is provided for diagnosing colon cancer in a sample suspected of being neoplastic. The method comprises the steps of:

comparing the level of at least one transcript in a first sample of a tissue to a second sample, wherein the first sample is of a colonic tissue suspected of being neoplastic and the second sample is of a normal human colonic tissue, and wherein the transcript is identified by a tag selected from the group consisting of those shown in Table 2;

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identifying the first sample as neoplastic when the level of the at least one transcript is found to be higher in the first sample than in the second sample.

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In another embodiment of the invention an isolated and purified human nucleic acid molecule is provided. The molecule comprises a SAGE tag selected from SEQ ID NO:1-732.

In yet another aspect of the invention an isolated nucleotide probe is provided. The probe comprises at least 12 nucleotides of a human nucleic acid molecule, wherein the human nucleic acid molecule comprises a SAGE tag selected from SEQ ID NO: 1-732.

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According to another aspect of the invention a method is provided for diagnosing pancreatic cancer in a sample suspected of being neoplastic. The method comprises the steps of:

comparing the level of at least one transcript in a first sample of a tissue to a second sample, wherein the first sample is of a pancreatic tissue suspected of being neoplastic and the second sample is of a normal human colon tissue, wherein said transcript is identified by a tag selected from the group consisting of those shown Table 4;

identifying the first sample as neoplastic when the level of the at least one transcript is found to be higher in the first sample than in the second sample.

According to still another embodiment of the invention a method of diagnosing cancer in a sample suspected of being neoplastic is provided. The method comprises the steps of:

comparing the level of at least one transcript in a first sample of a tissue to a second sample, wherein the first sample is of a tissue suspected of being neoplastic and the second sample is of a normal human tissue, wherein said transcript is identified by a tag selected from the group consisting of those shown Table 5;

identifying the first sample as neoplastic when the level of the at least one transcript is found to be higher in the first sample than in the second sample.

According to another embodiment of the invention a method is provided to aid in the determination of a prognosis for a colon cancer patient. The method comprises the steps of:

comparing the level of at least one transcript in a first sample of a tissue to a second sample, wherein the first sample is of a neoplastic colonic tissue and the second sample is of a normal human colonic tissue, and wherein the transcript is identified by a tag selected from the group consisting of those shown in Table 3; determining a poorer prognosis if the level of the at least one transcript is found to be lower in the first sample than in the second sample.

According to another aspect of the invention a method to aid in determining a prognosis for a patient with colon cancer is provided. The method comprises the steps of:

comparing the level of at least one transcript in a first tissue sample to a second sample, wherein the first sample is of a colonic cancer tissue and the second sample is of a normal human colonic tissue, and wherein the transcript is identified by a tag selected from the group consisting of those shown in Table 2;

determining a poorer prognosis if the level of the at least one transcript is found to be higher in the first sample than in the second sample.

In yet another embodiment of the invention a method is provided for diagnosing colon cancer in a sample suspected of being neoplastic. The method comprises the steps of:

comparing the level of expression of at least one protein in a first sample of a tissue to a second sample, wherein the first sample is of a colonic tissue suspected of being neoplastic and the second sample is of a normal human colonic tissue, and wherein the protein is encoded by a transcript identified by a tag selected from the group consisting of those shown in Table 3;

identifying the first sample as neoplastic when the level of expression of the protein is found to be lower in the first sample than in the second sample.

In another aspect of the invention a method of diagnosing colon cancer in a sample suspected of being neoplastic is provided. The method comprises the steps of:

comparing the level of expression of at least one protein in a first sample of a tissue to a second sample, wherein the first sample is of a colonic tissue suspected of being neoplastic and the second sample is of a normal human colonic tissue, and wherein the protein is encoded by a transcript

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identified by a tag selected from the group consisting of those shown in Table 2;

identifying the first sample as neoplastic when expression of the protein is found to be higher in the first sample than in the second sample.

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According to another embodiment of the invention a method is provided to aid in determining a prognosis of a patient having pancreatic cancer. The method comprises the steps of:

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comparing the level of at least one transcript in a first sample of a tissue to a second sample, wherein the first sample is of a neoplastic pancreatic tissue and the second sample is of a normal human colon tissue, wherein said transcript is identified by a tag selected from the group consisting of those shown Table 4;

determining a poorer prognosis if transcription is found to be higher in the first sample than in the second sample.

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In yet another aspect of the invention a method to aid in providing a prognosis for a cancer patient is provided. The method comprises the steps of:

comparing the level of at least one transcript in a first sample of a tissue to a second sample, wherein the first sample is of a neoplastic tissue and the second sample is of a normal human tissue of the same tissue type, wherein said transcript is identified by a tag selected from the group consisting of those shown Table 5;

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determining a poorer prognosis if transcription is found to be higher in the first sample than in the second sample.

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According to still another aspect of the invention, a method is provided for diagnosing pancreatic cancer in a sample suspected of being neoplastic. The method comprises the steps of:

comparing the level of expression of at least one protein encoded by a transcript in a first sample of a tissue to a second sample, wherein the first sample is of a pancreatic tissue suspected of being neoplastic and the second sample is of a normal human colon tissue, wherein said protein is

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encoded by a transcript identified by a tag selected from the group consisting of those shown Table 4;

identifying the first sample as neoplastic when expression of the protein is found to be higher in the first sample than in the second sample.

According to yet another aspect of the invention a method is provided for diagnosing cancer in a sample suspected of being neoplastic. The method comprises the steps of:

comparing the level of expression of at least one protein in a first sample of a tissue to a second sample, wherein the first sample is of a tissue suspected of being neoplastic and the second sample is of a normal human tissue, wherein said protein is encoded by a transcript identified by a tag selected from the group consisting of those shown Table 5;

identifying the first sample as neoplastic when expression of the protein is found to be higher in the first sample than in the second sample.

In still another embodiment of the invention a method is provided to aid in the determination of a prognosis of a colon cancer patient. The method comprises the steps of:

comparing the level of expression of at least one protein in a first sample of a tissue to a second sample, wherein the first sample is of a neoplastic colonic tissue and the second sample is of a normal human colonic tissue, and wherein the protein is encoded by a transcript identified by a tag selected from the group consisting of those shown in Table 3;

determining a poorer prognosis if the level of expression is found to be lower in the first sample than in the second sample.

In still another embodiment of the invention a method is provided to aid in determining a prognosis for a patient with colon cancer. The method comprises the steps of:

comparing the level of expression of at least one protein in a first tissue sample to a second sample, wherein the first sample is of a colonic cancer tissue and the second sample is of a normal human colonic tissue, and

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wherein the protein is encoded by a transcript identified by a tag selected from the group consisting of those shown in Table 2;

determining a poorer prognosis if the level of expression is found to be higher in the first sample than in the second sample.

In still another aspect of the invention a method is provided to aid in determining a prognosis of a patient having pancreatic cancer. The method comprises the steps of:

comparing the level of expression of at least one protein in a first sample of a tissue to a second sample, wherein the first sample is of a neoplastic pancreatic tissue and the second sample is of a normal human colon tissue, wherein said protein is encoded by a transcript identified by a tag selected from the group consisting of those shown Table 4;

determining a poorer prognosis if the level of expression is found to be higher in the first sample than in the second sample.

According to even a further aspect of the invention a method is provided to aid in providing a prognosis for a cancer patient. The method comprises the steps of:

comparing the level of expression of at least one protein in a first sample of a tissue to a second sample, wherein the first sample is of a neoplastic tissue and the second sample is of a normal human tissue of the same tissue type, wherein said protein is encoded by a transcript identified by a tag selected from the group consisting of those shown Table 5;

determining a poorer prognosis if the level of expression is found to be higher in the first sample than in the second sample.

In still another embodiment of the invention a method of treating a cancer cell is provided. The method comprises the step of:

administering to a cancer cell an antibody which specifically binds to a protein encoded by a transcript identified by a tag selected from the group consisting of those shown in Tables 2, 4, and 5, wherein the antibody is linked to a cytotoxic agent. In another aspect of the invention an antibody linked to a cytotoxic agent is provided. The antibody specifically binds to a protein encoded by a transcript identified by a tag selected from the group consisting of those shown in Tables 2, 4, and 5.

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According to another aspect of the invention, a method of detecting colon cancer in a patient is provided. The method comprises the steps of:

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comparing the level of at least one protein or transcript in a first body sample to a second body sample, wherein the first sample is a body sample of the patient and the second sample is of a normal human, wherein the protein is encoded by a transcript and the transcript is identified by a tag selected from the group consisting of those shown in Table 2, wherein the first and second body sample is a sample selected from the group consisting of blood, urine, feces, sputum, and serum;

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identifying neoplasia when the level of the at least one protein or transcript is found to be higher in the first sample than in the second sample.

In another aspect of the invention a method of detecting pancreatic cancer in a patient is provided. The method comprises the steps of:

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comparing the level of at least one protein or transcript encoded by a transcript in a first sample of a tissue to a second sample, wherein the first sample is of the patient and the second sample is of a normal human, wherein said protein is encoded by a transcript and the transcript is identified by a tag selected from the group consisting of those shown Table 4, wherein the first and second sample is a sample selected from the group consisting of blood, urine, feces, sputum, and serum;

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identifying neoplasia when the level of the at least one protein or transcript is found to be higher in the first sample than in the second sample.

Also provided by the present invention is a method of detecting cancer in a patient. The method comprises the steps of:

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comparing the level of at least one protein or transcript in a first sample to a second sample, wherein the first sample is of patient and the second sample is of a normal human, wherein said protein is encoded by a transcript and the transcript is identified by a tag selected from the group consisting of those shown Table 5, wherein the first and second body sample is a sample selected from the group consisting of blood, urine, feces, sputum, and serum;

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identifying neoplasia when the level of the at least one protein or transcript is found to be higher in the first sample than in the second sample.

Additionally provided by the present invention is a method to aid in the determination of a prognosis for a colon cancer patient. The method comprises the steps of:

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comparing the level of at least one protein or transcript in a first sample to a second sample, wherein the first sample is of a colon cancer patient and the second sample is of a normal human, wherein the protein is encoded by a transcript and the transcript is identified by a tag selected from the group consisting of those shown in Table 3, wherein the first and second body sample is a sample selected from the group consisting of blood, urine, feces, sputum, and serum;

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determining a poorer prognosis if the level of the at least one protein or transcript is found to be lower in the first sample than in the second sample.

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Provided by another embodiment of the invention is a method to aid in determining a prognosis for a patient with colon cancer. The method comprises the steps of:

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comparing the level of at least one protein or transcript in a first sample to a second sample, wherein the first sample is of a colonic cancer patient and the second sample is of a normal human, wherein the protein is encoded by a transcript and the transcript is identified by a tag selected from the group consisting of those shown in Table 2, wherein the first and second sample is a sample selected from the group consisting of blood, urine, feces, sputum, and serum;

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determining a poorer prognosis if the level of the at least one protein or transcript is found to be higher in the first sample than in the second sample.

According to still another aspect of the invention, a method to aid in determining a prognosis of a patient having pancreatic cancer is provided. The method comprises the steps of:

comparing the level of at least one protein or transcript in a first sample to a second sample, wherein the first sample is of a pancreatic cancer patient and the second sample is of a normal human, wherein said protein is encoded by a transcript and the transcript is identified by a tag selected from the group consisting of those shown Table 4, wherein said first and second sample is a sample selected from the group consisting of blood, urine, feces, sputum, and serum;

determining a poorer prognosis if the level of the at least one protein or transcript is found to be higher in the first sample than in the second sample.

Also provided by the present invention is a method to aid in providing a prognosis for a cancer patient. The method comprises the steps of

comparing the level of expression of at least one protein or transcript in a first sample to a second sample, wherein the first sample is of a cancer patient and the second sample is of a normal human, wherein said protein is encoded by a transcript and the transcript is identified by a tag selected from the group consisting of those shown Table 5, wherein the first and second sample is a sample selected from the group consisting of blood, urine, feces, sputum, and serum;

determining a poorer prognosis if the level of the at least one protein or transcript is found to be higher in the first sample than in the second sample.

The present invention further includes antisense oligonucleotides complementary in whole or in part to SEQ ID NOS:1-732.

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This invention also provides a method for screening for candidate agents that modulate the expression of a polynuleotide selected from the group consisting of the polynucleotides in SEQ ID NOS.1-732 or their respective complements, by contacting a test agent with a pancreatic or colon cell and monitoring expression of the polynucleotide, wherein the test agent which modifies the expression of the polynucleotide is a candidate agent.

The present invention provides the art with new methods and reagents for diagnosing and prognosing cancers. In addition, some of the newly disclosed genes may play an important role in the development of cancers.

BRIEF DESCRIPTION OF THE DRAWINGS

Fig. 1. Comparison of expression patterns in colorectal cancers and normal colon epithelium. (FIG. 1A) A semi-logarithmic plot reveals 51 tags that were decreased more than 10 fold in primary CR cancer cells whereas 32 tags were increased more than 10 fold. 62,168 and 60,878 tags derived from normal colon epithelium and primary CR cancers, respectively, were used for this analysis. The relative expression of each transcript was determined by dividing the number of tags observed in tumor and normal tissue as indicated. To avoid division by 0, a tag value of 1 was used for any tag that was not detectable in one of the samples. These ratios were then rounded to the nearest integer and their distribution plotted on the abscissa. The number of genes displaying each ratio was plotted on the ordinate. Tu: CR tumors; NC: Normal colon. (FIG. 1B and FIG. 1C) Differentially expressed genes in The number of transcripts found to be differentially colorectal cancers. expressed (P < 0.01) are presented as Venn diagrams. Diagrams of transcripts that were decreased (FIG. 1B) or increased (FIG. 1C) in CR cancers compared to normal colon epithelium. Comparisons were between primary tumors and cells in culture as indicated.

Fig. 2. Northern blot analysis of genes differentially expressed in gastrointestinal neoplasia. Northern blot analysis was performed on total RNA (5 μg isolated from primary CR carcinomas (T) and matching normal colon epithelium (N), or pancreatic carcinomas. The top panel in each case show an

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example of the ethidium bromide stained gels prior to transfer. The number of SAGE tags observed in the original analysis is indicated to the right of each blot. (FIG. 2A) Examples of transcripts that were decreased or increased in CR cancers. (FIG.2B) Examples of transcripts increased in pancreatic cancers (10). (FIG.2C) Examples of transcripts elevated in cancer which were or were not cancer type specific. Probes used for Northern blot analysis were as follows (Human SAGE Tag unique identifier, gene name, (GenBank accession number)): (FIG. 2A) H204104, Guanylin (M95714); H259108, (see Table 2); H1000193, (see Table 2); H998030, (see Table 2). (FIG. 2B) H294155, RIG-E (U42376); H560056, TIMP-1 (S68252). (FIG. 2C) H802810, EST338411 (W52120); H85882, 1-8D (X57351); H618841, GA733-1 (X13425).

Tables 2-5. Transcripts Differentially Expressed in Human Cancer.

Tag sequence represents the NlaIII site plus the adjacent 11 bp SAGE tag. Tag number indicates a SAGE UID (unique identifier). NC, TU, CL, PT, PC, refers to the number of the indicated tag observed in RNA isolated from normal colorectal epithelium, primary colorectal cancers, colorectal cancer cell lines, primary pancreatic cancers, or pancreatic cancer cell lines, respectively. The Accession and Gene Name refer to representative GenBank entries that contain the tag sequence.

Table 2 Transcripts increased in colorectal cancer.

Table 3 Transcripts decreased in colorectal cancer.

Table 4 Transcripts increased in pancreatic cancer.

Table 5 Transcripts increased in pancreatic and colorectal cancer.

25 <u>DETAILED DESCRIPTION</u>

The inventors have discovered sets of human genes which are either upregulated or downregulated in cancer cells, as compared to normal cells. Specifically, certain genes have been found to be upregulated or downregulated in colorectal and/or pancreatic cancer cells, when compared to normal colon

cells. These sets of differentially regulated genes can be used as diagnostic markers, either individually or in sets of, for example, 2, 5, 10, 20, or 30.

Genes whose expression was detected to be increased in colorectal cancer are shown in Table 2. Genes whose expression was detected to be decreased in colorectal cancer are shown in Table 3. Genes whose expression was detected as increased in pancreatic cancer are shown in Table 4. Genes whose expression was detected as increased in both pancreatic cancer and colorectal cancer are shown in Table 5. These latter genes likely play a role in neoplastic development generally.

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Tag sequences, as provided herein, uniquely identify genes. This is due to their length, and their specific location (3') in a gene from which they are drawn. The full length genes can be identified by matching the tag to a gene data base member, or by using the tag sequences as probes to physically isolate previously unidentified genes from cDNA libraries. The methods by which genes are isolated from libraries using DNA probes are well known in the art. See, for example, Veculescu et al., Science 270: 484 (1995), and Sambrook et al. (1989), MOLECULAR CLONING: A LABORATORY MANUAL, 2nd ed. (Cold Spring Harbor Press, Cold Spring Harbor, New York). Once a gene or transcript has been identified, either by matching to a data base entry, or by physically hybridizing to a cDNA molecule, the position of the hybridizing or matching region in the transcript can be determined. If the tag sequence is not in the 3' end, immediately adjacent to the restriction enzyme used to generate the SAGE tags, then a spurious match may have been made. Confirmation of the identity of a SAGE tag can be made by comparing transcription levels of the tag to that of the identified gene in certain cell types.

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In addition to the sequences shown in SEQ ID NOS: 1-732, or their complements, this invention also provides the anti-sense polynucleotide stand, e.g. antisense RNA to these sequences or their complements. One can obtain an antisense RNA using the sequences provided in SEQ ID NOS: 1-732 and the methodology described in Vander Krol et al. (1988) BioTechniques 6:958.

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The invention also encompasses polynucleotides which differ from that of the polynucleotides described above, but which produce the same phenotypic effect, such as the allele. These altered, but phenotypically equivalent polynucleotides are referred to "equivalent nucleic acids." This invention also encompasses polynucleotides characterized by changes in non-coding regions that do not alter the phenotype of the polypeptide produced therefrom when compared to the polynucleotide herein. This invention further encompasses polynucleotides, which hybridize to the polynucleotides of the subject invention under conditions of moderate or high stringency.

The polynucleotides can be conjugated to a detectable marker, e.g., an

enzymatic label or a radioisotope for detection of nucleic acid and/or expression of the gene in a cell. A wide variety of appropriate detectable markers are known in the art, including fluorescent, radioactive, enzymatic or other ligands, such as avidin/biotin, which are capable of giving a detectable signal. In preferred embodiments, one will likely desire to employ a fluorescent label or an enzyme tag, such as urease, alkaline phosphatase or peroxidase, instead of radioactive or other environmental undesirable reagents. In the case of enzyme tags, colorimetric indicator substrates are known which can be employed to provide a means visible to the human eye or spectrophotometrically, to identify specific hybridization with complementary nucleic acid-containing samples. Briefly, this invention further provides a method for detecting a single-stranded polynucleotide identified by SEO ID NOS.1-732 or its complement, by contacting target single-stranded polynucleotides with a labeled, single-stranded polynucleotide (a probe) which is at least 10 nucleotides of the complement of SEQ ID NOS: 1-732 (or the corresponding complement) under conditions permitting hybridization (preferably moderately stringent hybridization conditions) of complementary

single-stranded polynucleotides, or more preferably, under highly stringent

hybridization conditions. Hybridized polynucleotide pairs are separated from un-hybridized, single-stranded polynucleotides. The hybridized polynucleotide

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pairs are detected using methods well known to those of skill in the art and set forth, for example, in Sambrook et al. (1989) supra.

The polynucleotides of this invention can be isolated using the technique described in the experimental section or replicated using PCR. The PCR technology is the subject matter of United States Patent Nos. 4.683, 195. 4,800,159, 4,754,065, and 4,683,202 and described in PCR: The Polymerase Chain Reaction (Mullis et al. eds, Birkhauser Press, Boston (1994)) or MacPherson et al. (1991) and (1994), supra, and references cited therein. Alternatively, one of skill in the art can use the sequences provided herein and a commercial DNA synthesizer to replicate the DNA. Accordingly, this invention also provides a process for obtaining the polynucleotides of this invention by providing the linear sequence of the polynucleotide, nucleotides, appropriate primer molecules, chemicals such as enzymes and instructions for their replication and chemically replicating or linking the nucleotides in the proper orientation to obtain the polynucleotides. In a separate embodiment, these polynucleotides are further isolated. Still further, one of skill in the art can insert the polynucleotide into a suitable replication vector and insert the vector into a suitable host cell (procarvotic or eucarvotic) for replication and amplification. The DNA so amplified can be isolated from the cell by methods well known to those of skill in the art. A process for obtaining polynucleotides by this method is further provided herein as well as the polynucleotides so obtained.

RNA can be obtained by first inserting a DNA polynucleotide into a suitable host cell. The DNA can be inserted by any appropriate method, e.g., by the use of an appropriate gene delivery vector or by electroporation. When the cell replicates and the DNA is transcribed into RNA; the RNA can then be isolated using methods well known to those of skill in the art, for example, as set forth in Sambrook et al. (1989) supra. For instance, mRNA can be isolated using various lytic enzymes or chemical solutions according to the procedures set forth in Sambrook et al. (1989), supra or extracted by nucleic-acid-binding resins following the accompanying instructions provided by manufactures.

Polynucleotides having at least 10 nucleotides and exhibiting sequence complementarity or homology to SEQ ID NOS: 1-732 find utility as hybridization probes. In some aspects, the full coding sequence of the transcript, i.e., for SEQ ID NOS: 1-732, are known. Accordingly, any portion of the known sequences available in GenBank, or homologous sequences, can be used in the methods of this invention.

It is known in the art that a "perfectly matched" probe is not needed for a specific hybridization. Minor changes in probe sequence achieved by substitution, deletion or insertion of a small number of bases do not affect the hybridization specificity. In general, as much as 20% base-pair mismatch (when optimally aligned) can be tolerated. Preferably, a probe useful for detecting the aforementioned mRNA is at least about 80% identical to the homologous region of comparable size contained in the previously identified sequences identified by SEQ ID NOS:1-732, which correspond to previously characterized genes or SEQ ID NOS:1-732, which correspond to known ESTs. More preferably, the probe is 85% identical to the corresponding gene sequence after alignment of the homologous region; even more preferably, it exhibits 90% identity.

These probes can be used in radioassays (e.g. Southern and Northern blot analysis) to detect, prognose, diagnose or monitor various pancreatic or colon cells or tissue containing these cells. The probes also can be attached to a solid support or an array such as a chip for use in high throughput screening assays for the detection of expression of the gene corresponding to one or more polynucleotide(s) of this invention. Accordingly, this invention also provides at least one of the transcripts identified as SEQ ID NOS:1-732, or its complement, attached to a solid support for use in high throughput screens.

The total size of fragment, as well as the size of the complementary stretches, will depend on the intended use or application of the particular nucleic acid segment. Smaller fragments will generally find use in hybridization embodiments, wherein the length of the complementary region may be varied,

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such as between about 10 and about 100 nucleotides, or even full length according to the complementary sequences one wishes to detect.

Nucleotide probes having complementary sequences over stretches greater than 10 nucleotides in length are generally preferred, so as to increase stability and selectivity of the hybrid, and thereby improving the specificity of particular hybrid molecules obtained. More preferably, one can design polynucleotides having gene-complementary stretches of more than 50 nucleotides in length, or even longer where desired. Such fragments may be readily prepared by, for example, directly synthesizing the fragment by chemical means, by application of nucleic acid reproduction technology, such as the PCR technology with two priming oligonucleotides as described in U.S. Pat. No. 4,603,102 or by introducing selected sequences into recombinant vectors for recombinant production. A preferred probe is about 50-75 or more preferably, 50-100, nucleotides in length.

The polynucleotides of the present invention can serve as primers for the detection of genes or gene transcripts that are expressed in pancreatic or colon cells. In this context, amplification means any method employing a primer-dependent polymerase capable of replicating a target sequence with reasonable fidelity. Amplification may be carried out by natural or recombinant DNA-polymerases such as T7 DNA polymerase, Klenow fragment of E.coli DNA polymerase, and reverse transcriptase.

A preferred amplification method is PCR. However, PCR conditions used for each reaction are empirically determined. A number of parameters influence the success of a reaction. Among them are annealing temperature and time, extension time, Mg²⁺ ATP concentration, pH, and the relative concentration of primers, templates, and deoxyribonucleotides. After amplification, the resulting DNA fragments can be detected by agarose gel electrophoresis followed by visualization with ethidium bromide staining and ultraviolet illumination.

The invention further provides the isolated polynucleotide operatively linked to a promoter of RNA transcription, as well as other regulatory

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sequences for replication and/or transient or stable expression of the DNA or RNA. As used herein, the term "operatively linked" means positioned in such a manner that the promoter will direct transcription of RNA off the DNA molecule. Examples of such promoters are SP6, T4 and T7. In certain embodiments, cell-specific promoters are used for cell-specific expression of the inserted polynucleotide. Vectors which contain a promoter or a promoter/enhancer, with termination codons and selectable marker sequences, as well as a cloning site into which an inserted piece of DNA can be operatively linked to that promoter are well known in the art and commercially available. For general methodology and cloning strategies, see Gene Expression Technology (Goeddel ed., Academic Press, Inc. (1991)) and references cited therein and Vectors: Essential Data Series (Gacesa and Ramji, eds., John Wiley & Sons, N.Y. (1994)), which contains maps, functional properties, commercial suppliers and a reference to GenEMBL accession numbers for various suitable vectors. Preferable, these vectors are capable of transcribing RNA in vitro or in vivo.

Fragment of the sequences shown in SEQ ID NOS:1-732 or their respective complements also are encompassed by this invention, preferably at least 10 nucleotides and more preferably having at least 18 nucleotides. Larger polynucleotides, e.g., cDNA or genomic DNA, which hybridize under moderate or stringent conditions to the polynucleotide sequences shown in SEQ ID NOS:1-732, or their respective complements, also are encompassed by this invention.

In one embodiment, these fragments are polynucleotides that encode polypeptides or proteins having diagnostic and therapeutic utilities as described herein as well as probes to identify transcripts of the protein which may or may not be present. These nucleic acid fragments can by prepared, for example, by restriction enzyme digestion of the polynucleotide of SEQ ID NOS:1-732, or their complements, and then labeled with a detectable marker. Alternatively, random fragments can be generated using nick translation of the molecule. For

methodology for the preparation and labeling of such fragments, see Sambrook et al., (1989) supra.

Expression vectors containing these nucleic acids are useful to obtain host vector systems to produce proteins and polypeptides. It is implied that these expression vectors must be replicable in the host organisms either as episomes or as an integral part of the chromosomal DNA. Suitable expression vectors include viral vectors, including adenoviruses, adeno-associated viruses, retroviruses, cosmids, etc. Adenoviral vectors are particularly useful for introducing genes into tissues in vivo because of their high levels of expression and efficient transformation of cells both in vitro and in vivo. When a nucleic acid is inserted into a suitable host cell, e.g., a procaryotic or a eucaryotic cell and the host cell replicates, the protein can be recombinantly produced. Suitable host cells will depend on the vector and can include mammalian cells, animal cells, human cells, simian cells, insect cells, yeast cells, and bacterial cells constructed using well known methods. See Sambrook et al. (1989) supra. In addition to the use of viral vector for insertion of exogenous nucleic acid into cells, the nucleic acid can be inserted into the host cell by methods well known in the art such as transformation for bacterial cells; transfection using calcium phosphate precipitation for mammalian cells; or DEAE-dextran; electroporation; or microinjection. See Sambrook et al. (1989) supra for this methodology. Thus, this invention also provides a host cell, e.g. a mammalian cell, an animal cell (rat or mouse), a human cell, or a procaryotic cell such as a bacterial cell, containing a polynucleotide encoding a protein or polypeptide or antibody.

When the vectors are used for gene therapy in vivo or ex vivo, a pharmaceutically acceptable vector is preferred, such as a replication-incompetent retroviral or adenoviral vector. Pharmaceutically acceptable vectors containing the nucleic acids of this invention can be further modified for transient or stable expression of the inserted polynucleotide. As used herein, the term "pharmaceutically acceptable vector" includes, but is not limited to, a vector or delivery vehicle having the ability to selectively target

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and introduce the nucleic acid into dividing cells. An example of such a vector is a "replication-incompetent" vector defined by its inability to produce viral proteins, precluding spread of the vector in the infected host cell. An example of a replication-incompetent retroviral vector is LNL6 (Miller, A.D. et al. (1989) BioTechniques 7:980-990). The methodology of using replication-incompetent retroviruses for retroviral-mediated gene transfer of gene markers is well established (Correll et al. (1989) PNAS USA 86:8912; Bordignon (1989) PNAS USA 86:8912-52; Culver, K. (1991) PNAS USA 88:3155; and Rill, D.R. (1991) Blood 79(10):2694-700. Clinical investigations have shown that there are few or no adverse effects associated with the viral vectors, see Anderson (1992) Science 256:808-13.

Compositions containing the polynucleotides of this invention, in isolated form or contained within a vector or host cell are further provided herein. When these compositions are to be used pharmaceutically, they are combined with a pharmaceutically acceptable carrier.

This invention further encompasses genes, either genomic or cDNA, which code for a polypeptide or protein in the cell of interest. The genes specifically hybridize under moderate or stringent conditions to a polynucleotide identified by SEQ ID NOS: 1-732 or their respective complements. The process of identification of larger fragment or the full-length coding sequence to which the partial sequence depicted in SEQ ID NOS:1-732 hybridizes preferably involves the use of the methods and reagents provided in this invention, either singularly or in combination.

Five methods are disclosed herein which allows one of skill in the art to isolate the gene or cDNA corresponding to the transcripts of the invention.

RACE-PCR Technique

One method to isolate the gene or cDNA which code for a polypeptide or protein and which corresponds to a transcript of this invention, involves the 5'-RACE-PCR technique. In this technique, the poly-A mRNA that contains the coding sequence of particular interest is first identified by hybridization to

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a sequence disclosed herein and then reverse transcribed with a 3'-primer comprising the sequence disclosed herein. The newly synthesized cDNA strand is then tagged with an anchor primer of a known sequence, which preferably contains a convenient cloning restriction site attached at the 5'end. The tagged cDNA is then amplified with the 3'-primer (or a nested primer sharing sequence homology to the internal sequences of the coding region) and the 5'-anchor primer. The amplification may be conducted under conditions of various levels of stringency to optimize the amplification specificity. 5'-RACE-PCR can be readily performed using commercial kits (available from, e.g., BRL Life Technologies Inc, Clotech) according to the manufacturer's instructions.

Identification of known genes or ESTs

In addition, databases exist that reduce the complexity of ESTs by assembling contiguous EST sequences into tentative genes. For example, TIGR has assembled human ESTs into a datable called THC for tentative human consensus sequences. The THC database allows for a more definitive assignment compared to ESTs alone. Software programs exist (give examples) that allow for assembling ESTs into contiguous sequences from any organism.

Isolation of cDNAs from a library by probing with the SAGE transcript or tag

Alternatively, mRNA from a sample preparation was used to construct cDNA library in the ZAP Express vector following the procedure described in Velculescu et al. (1997) Science 270:484. The ZAP Express cDNA synthesis kit (Stratagene) was used accordingly to the manufacturer's protocol. Plates containing 250 to 2000 plaques are hybridized as described in Rupert et al. (1988) Mol. Cell. Bio. 8:3104 to oligonucleotide probes with the same conditions previously described for standard probes except that the hybridization temperature is reduced to room temperature. Washes are performed in 6X standard-saline-citrate 0.1% SDS for 30 minutes at room temperature. The probes are labeled with 32P-ATP through use of T4 polynucletoide kinase.

Table 2 - Transcripts increased in colon cancer

Transcripts increased in only colon primary tumors compared to normal colon (61 genes)

#	Tag Sequence	Tag Number	S	TO	CF	PT	PC	Accession	Сепе Мате
1	CATGCACCTAATTGG	H285759	612	755	411	161	333	F15516	H.sapiens mitochondrial EST sequence (1-t-12) from
2	CATGTGATTTCACTT	H933704	452	595	235	80	314	U35430	Human cytochrome c oxidase subunit III (COIII) pse
~	CATGCCTGTAATCCC	H388150	433	249	380	443	161	Z70701	H.sapiens mRNA (fetal brain cDNA c2_11).
								X71347	H.sapiens HNF1-C mRNA.
						-		X71346	H.sapiens HNF1-B mRNA.
4	CATGCACTACTCACC	H291282	293	527	78	14	83	U09500	Human mitochondrion cytochrome b gene, partial cds
0	S CATGGTGAAACCCCA(G)	H753750	392	517	389	453	194	X66785	H.sapiens mRNA for transacylase (DBT).
								X17648	Human mRNA for granulocyte-macrophage colony-stimu
								U09087	Human thymopoietin beta mRNA, complete cds.
								U09088	Human thymopoietin gamma mRNA, complete cds.
								U20770	Human metastasis suppressor (KAII) mRNA, complete
9	CATGGGCTTTAGGGA	H687915	37	372	9	29	=	W15552	2b91h11.s1 Soares parathyroid tumor NbHPA Homo sap
								W32091	zc05d03.s1 Soares paraliyroid tumor Nbl·IPA Homo sap
								R62866	yi 11d07,r1 Homo sapiens cDNA clone 138925 5'.
7	CATGACTTTCCAAA	H130369	32	272	32	23	20	X89839	H.sapiens mitochondrial DNA for loop attachment se
8	CATGTGGTGTATGCA	H965434	23	171	9	30	2	T11555	A 1486F Homo sapiens cDNA clone A 1486 similar to Mi
6	CATGAGGGTGTTTTC	H175872	26	218	7	20	10	T15773	IB1870 Homo sapiens cDNA 3'end similar to Human mi
9	CATGAGGTCAGGAGA(T)	H177315	93	213	113	148	58	X12544	Human mRNA for HLA class II DR-beta (HLA-DR B).
								S73483	phosphorylase kinase catalytic subunit PHKG2 homol
=	CATGTTGGCCAGGCT	H1025322	124	194	63	111	51	X74301	H.sapiens mRNA for MHC class II transactivator.
								U28687	Human zinc finger containing protein ZNF157 (ZNF15
								U29119	Human leiomyoma LM-196.4 ectopic sequence from HMG
								U\$6236	Human Fc alpha receptor b mRNA, complete cds.
2	CATGATCACGCCCTC	H214616	97	186	17	41	49	W03751	za62h11.rl Soares fetal liver spleen INFLS Homo sa
								W03770	za63f10.r1 Soares fetal liver spleen INFLS Homo sa

							W04748	224200 rl Spares fetal liver spleep INFLS Homo sa
ייייייייייייייייייייייייייייייייייייייי	169669H	37	2	=	191	6	T12078	A730R Homo sapiens cDNA clone A730 similar to Mito
CA10000167							W45641	zc26a12.s1 Soares senescent fibroblasts NbHSF Homo
CATCCCTAGGTTTAT	H641789	38	144	13	25	2	DS1017	Human fetal brain cDNA 3'-end GEN-007C04.
14 CA10001A0							D53694	Human fetal brain cDNA 3'-end GEN-117E01.
15 CATGCCCGTACATC	H350996	56	132	35	0	81		Unknown
15 CATGAGTAGGTGGC	H183018	18	3	7	=	-	D51021	Human fetal brain cDNA 3'-end GEN-007D07.
CALCACITY OF THE PROPERTY OF T							D51052	Human fetal brain cDNA 3'-end GEN-009C05.
							D52836	Human fetal brain cDNA 3'-end GEN-089E01.
17 CATGCCTGTAGTCCC	H388278	79	124	19	71	23	D83195	Human DNA for Deoxyribonuclease I precursor.
CATGAGACCCACAC	H136465	64	121	78	24	15	D54113	Human fetal brain cDNA 5'-end GEN-129B05.
CATGCATTTGTAATA	H327364	49	107	35	7	Q	F15796	H.sapiens mitochondrial EST sequence (102-25) from
20 CATGTCCCGTACCT	H874182	28	78	14	0	23		
21 CATGGCCAACCTCCT	H606582	23	73	8	9	6	Z59183	H.sapiens CpG island DNA genomic Msel fragment, cl
							D52905	Human fetal brain cDNA 5'-end GEN-091D11.
22 CATGGCGATCCCTT	H609624	29	73	7	14	91	F16449	H.sapiens mitochondrial EST sequence (129-09) from
CATGTTGGTCAGGCT	H1027370	35	29	18	35	14	U06452	Human melanoma antigen recognized by T-cells (MART
24 CATGTCCTATTAAG	H881603	20	49	17	15	82		
CATGITACTIATACT	H991026	2	47	2	_	4	D51004	Human fetal brain cDNA 3'-end GEN-006D02.
							L49057	Homo sapiens retinal fovea EST HFD010904 sequence.
							D51071	Human fetal brain cDNA 3'-end GEN-010E01.
CATGATGGCAGGAGT	HZ38755	=	45	-	4	2		
CATGCTAAGGCGAGG	H461411	5	44	2	3	3		
28 CATGGGTGAGACACT	H713234	7	44	20	13	15	J03592	Human ADP/ATP translocase mRNA, 3' end, clone pHAT
29 CATGACCTGTATCCC	H97078	9	42	17	100	32	X57352	Human I-8U gene from interferon-inducible gene fam
IN CATGCCAGTCCGCCT	H339302	0	39	0	_	0	H01571	yj33e06.rl Homo sapiens cDNA clone 150562 5' simil
							H03072	yj46g12.r1 Homo sapiens cDNA clone 151846 5' simil
11 CATCTAATTTTGCC	H802810	_	37	0	-	0	T25155	EST730 Homo sapiens cDNA clone 34C11.
12 CATGITTAGCITGITT	H993264	9	37	7	-	5	D50972	Human fetal brain cDNA 3'-end GEN-004A05.
							D51211	Human fetal brain cDNA 3'-end GEN-017E08.
					·		D52162	Human fetal brain cDNA 3'-end GEN-069F04.
		_					T23865	seq2012 Homo sapiens cDNA clone Cot1374Ft-4HB3MA-3
33 CATGGCCACCCCTG	H607576	0	35	1	0	0	M32053	Human H19 RNA gene, complete cds.
34 CATGTAATAAAGGTG	H798764	13	35	19	33	2	X67247	H.sapiens rpS8 gene for ribosomal protein S8.
15 CATGTACTGCTCGGA	H817627	13	32	\$	_	14	T11939	A953F Homo sapiens cDNA clone A953 similar to Mito
						i		

145	16 CATGGTGAAACCCA	H753749	6	31	22	30	4	T95857	ye42f01.s1 Homo sapiens cDNA clone 120409 3' simil
								W03237	za35b09.r1 Soares fetal liver spleen INFLS Homo sa
_								W03326	za63g03.rl Soares fetal liver spleen INFLS Homo sa
=	CATGGAAACTGAACA	H526210	9	26	11	5	3	X54195	Human line-1 element DNA, host sequence flanking t
							_	U29607	Human methionine aminopeptidase mRNA, complete cds
								H95100	yw57b10.rl Homo sapiens cDNA clone 256315 5' simil
82	CATGACTTTTAAAA	H131009	-	22	4	-	0		
2	19 CATGGACTGCGTGCC	HSSS450 "	0	21	7	6	[2]	D29062	Human keratinocyte cDNA, clone 067.
								D29563	Human keratinocyte cDNA, clone 713.
04	CATGTCAGTGGTAGT	H863923	4	71	7	2	-	T03196	FB3B5 Homo sapiens cDNA clone FB3B5 3'end.
2	CATGAAACTGTGGTT	H7916	2	20	7	7	-	257093	H.sapiens CpG DNA, clone 164a10, reverse read cpg1
:						_		Z60184	H.sapiens CpG island DNA genomic Mse1 fragment, cl
						_		Z63649	H.sapiens CpG island DNA genomic Msel fragment, cl
								W31349	zb95d06.s1 Soares parathyroid tumor NbHPA. Homo sap
42	CATGGGGGGGGGGGT	H699051	0	61	0	0	0		
12	41 CATGOTGCCGTGCC		7	6	-	0	0	W31448	zb96h01.si Soares parathyroid tumor NbHPA Homo sap
								W47282	zc40b06.rl Soares senescent fibroblasts NbHSF Homo
74	14 CATGGGGGTAACTA	11699144	3	6	2	12	2	X71428	H.sapiens fus mRNA.
							-	S62140	TLS=translocated in liposarcoma [human, mRNA, 1824
								W31782	zb96a06.r1 Soares parathyroid tumor NbHPA Homo sap
45	45 CATGTCCTGCCCCAT	H883029	3	16	14	77	91	M24398	Human parathymosin mRNA, complete cds.
46	CATGAAGTGGCAAGA	H47683	0	91	0	0	0		
5		H708358	0	91	0	0	0	U33317	Human defensin 6 (HD-6) gene, complete cds.
1								M98331	Homo sapiens defensin 6 mRNA, complete cds.
24	CATGGGCTACACCTT	H684312	2	16	0	2		D32027	Human mRNA for T cell receptor V beta 14 CDR3, par
1			2	9	0	2	_	T11701	A1225F Homo sapiens cDNA clone A1225 similar to Mi
49	49 CATGAGGGTGTTTCC	HI75870	_	15	0	0	0	D51783	Human fetal brain cDNA 5'-end GEN-051G02.
20	50 CATGCAAGGACCAGC	H272467	0	13	0	2		D13138	Human mRNA for dipeptidase.
							_		Homo sapiens (clones MDP4, MDP7) microsomal dipept
									RDP=renal dipeptidase [human, kidney, Genomic, 357
=	CATGTGGAAATGACC	H950498	0	2	٥	167	0	M10629	Human alpha-I collagen gene, 3' end with polyA sit
	CATGATCCGCCTGCC	H219514	-	13	3	4	_	H11641	ym17e04.s1 Homo sapiens cDNA clone 47962 3' simila
1								R95667	yq51a09.s1 Homo sapiens cDNA clone 199288 3' simil
2	S3 CATGTCCCGTACAC	H875282	-	13	0	0	_		
2	54 CATGATGTAAAAAT	H241665	0	=	0	12	4	M74090	Human TB2 gene mRNA, 3' end.

							103801	Human lysozyme mRNA, complete cds with an Alu repe
							M19045	M19045 Human lysozyme mRNA, complete cds.
	H337244	0	=	0	0	0		
	H85882	0	10	-	26	3	X57351	Human 1-8D gene from interferon-inducible gene fam
Ì							X02490	Human interferon-inducible mRNA (cDNA 1-8).
١	H165175	0	2	0	٥	0		
l	H243747	0	10	0	165	0	103040	J03040 Human SPARC/osteonectin mRNA, complete cds.
	H310975	0	10	9	7	4	U55217	U55217 Human RNA fragment from patients with Crohn's dise
l	H613862	0	10	2	15	7		
1	H992010	0	10	3	3	9	M94083	M94083 Human chaperonin-like protein (HTR3) mRNA, complet
l							L27706	L27706 Human chaperonin protein (Tcp20) gene complete cds
l								

Transcripts increased in both colon primary tumors and colon cancer cell lines compared to normal colon (47 genes)

NC: Normal Colon
TU: Colon Primary Turnor
CL: Colon Cancer Cell Line
PT: Pancreatic Primary Turnor
PC: Pancreatic Cancer Cell Line

**	Tag Sequence	Tag Number NC	N	IJ	C	77	PC	Accession	Gene Name
1-	CATGGCAGCCATCCG	H599350	87	180	230	72	138	U14969	Human ribosomal protein L28 mRNA, complete cds.
7	CATGATGGCTGGTAT	H239533	52	153	318	80	294	X17206	Human mRNA for LLRep3.
m	CATGCCCGTCCGGAA	H355689	87	142	246	178	250	X64707	H.sapiens BBCI mRNA
7	CATGAGGCTACGGAA	H171113	4	117	167	98	147	X56932	H.sapiens mRNA for 23 kD highly basic protein
100	CATGAGCACCTCCAG	H148949	42	116	161	103	190	Z11692	H.sapiens mRNA for elongation factor 2.
9	CATGCTGGGTTAATA	H502724	29	115	160	75	34	M81757	H.sapiens S19 ribosomal protein mRNA, complete cds
1	CATGGGATTTGGCCT	H671654	55	108	222	73	185	M17887	Human acidic ribosomal phosphoprotein P2 mRNA, com
∞	CATGTACCATCAATA	H807748	46	107	86	42	189	X53778	H.sapiens ling mRNA for uracil DNA glycosylase.
								102642	Human glyceraidehyde 3-phosphate dehydrogenase mRN
0	CATGTGGGCAAAGCC	H959498	51	103	156	45	152	Z11531	H.sapiens mRNA for elongation factor-1-gamma.
								M55409	Human pancreatic tumor-related protein mRNA, 3' en
2	CATGAATCCTGTGGA	H55227	30	95	102	48	156	Z28407	H.sapiens mRNA for ribosomal protein L8.
=	11 CATGGGACCACTGAA	H660601	36	- 92	114	43	63	X73460	H.sapiens mRNA for ribosomal protein L3.
12	12 CATGAGGGCTTCCAA	H174037	47	91	167	16	155	M73791	Human novel gene mRNA, complete cds.
		r						M64241	Human Wilm's tumor-related protein (QM) mRNA, comp
								835960	laminin receptor homolog (3' region) [human, mRNA
=	CATGAAGGTGGAGGA	H44683	48	91	182	13	215	X80822	H.sapiens mRNA for ORF.
14	14 CATGTGCACGTTITC	H935680	45	87	105	61	122	X03342	Human mRNA for ribosomal protein L32
15	15 CATGTCAGATCTTTG	H861056	37	81	93	50	8	M58458	Human ribosomal protein S4 (RPS4X) isoform mRNA, с
								M22146	Human scar protein mRNA, complete cds.
9	16 CATGTGGTGTTGAGG	H965603	42	79	83	55	250	X69150	H.sapiens mRNA for ribosomal protein S18.
								L06432	Homo sapiens 18S ribosomal protein (HKE3) mRNA seq
11	17 CATGCCTAGCTGGAT	H379369	28	77	80	46	143	Y00052	Human mRNA for T-cell cyclophilin.
==	18 CATGCTTGGGTTTTG	518912	0	73	42	0			Human DNA for insulin-like growth factor II (IGF-2);
2	19 CATGCTCCTCACCTG	H482584	12	72	4	*	S	U16811	Human Bak mRNA, complete cds.

	П				27 H.sapiens mRNA for ribosomal protein L19.	94 Human MHC protein homologous to chicken B complex								П				32 Human ribosomal protein L23a mRNA, partial cds.	70 Human ribosomal protein S5 mRNA, complete cds.						_				7					18 H.sapiens DNA for orphan TCR V-beta segment (allel
D14530	X73974	D23661	1.06505	M17886	X63527	M24194	U14967	X55954	X52839	H38868	H71935	Z43914	T48545	X04347	X00910	X61156	103799	U02032	U14970	59685X	M36981	L16785	L10376	\$80520	M77349	X58536	X00497	X16934	Y00345	X81005	D28137		W46476	X72718
100	8	119	118	66	146	77	19	120		ঞ				8	0	27		55	8	38			31		∞	8.	01	77	ম	3		_	9	Ц
48	55	55	42	49	77	35	2	49		21				ន	0	13		8	25	25			27		22	44	141	∞	=	Ξ			∞	Ц
116	183	73	82	154	17	78	20	5		B				ž	2	≅		8	2	25			8		15	11	25	37	27	91			15	
65	62	99	57	26	35	55	53	50		49				44	42	4		39	8	29			53		24	22	22	81	16	15			7	
17	28	6	15	12	24	2	82	23		15				6	0	14		0	12	2		L	6		0	0	3	7	_	_	L	_	0	
H507577	H416261	H274492	H79065	H1000193	H528694	H998030		H253260		H119809				H507455	802871	H524524		H33331	H390692	H125661			H302367		H769020	H760291	H774461	H918273	H2056	H948604			H495251	
20 10 ATCOTOTTGGTGAT	21 CATGCGCCGGAACAC	CATGCAATAAA	CATGACATCAT	24 CATGITCAATAAAA	25 CATOCA ACACATICA	25 CATGTTATGGGATGT	22 CATGGCATAATAGGT		28 CA10A11A11A1	20 CATGACTCCAAAAA	מאסוסעסו ער			30 CATGCTGTTGATTGC	21 CATGTACAAAATCGA	CATGGAAAAT	CATOCACAC	13 CATCAAGAAGATAGA	CATGOOTTOGAC	35 CATGACTGGGTCTAT			36 CATGCAGCTCACTGA		37 CATGGTGTGTTGTA	CATGGTGCGCT				Т.	2012010100		42 CATGCTGATGGCAGA	מיומרומיו

									Soares fetal heart NbHH19W Homo sapiens cDNA clone 342926
7	4.1 CATGACTCGCTCTGT	H121311	0	12	91	5	7	0 12 16 5 7 H121311	3'.
									EST176663 Colon carcinoma (Caco-2) cell line II Homo sapiens
								AA305589	AA305589 cDNA 5' end
\$	45 CATGGCCCAAGGACC	4	0	12	16	82	17	X53416	H610466 0 12 19 82 17 X53416 Human mRNA for actin-binding protein (filamin) (AB
46	46 CATGATCTTGTTACT		0	П	28	<i>L</i> 9	0	X02761	X02761 Human mRNA for fibronectin (FN precursor).
47	47 CATGAAGCTGCTGGA		0	10	17	9	9	Z26305	H40571 0 10 17 6 6 226305 H.sapiens isoform I gene for L-type calcium channe

20 CATGCCCTGGGTTCT

cell lines compared to normal colon (181 genes) Transcripts increased in only colon cancer

NC: Normal Colon

TU: Colon Primary Tumor CL: Colon Cancer Cell Line PT: Pancreatic Primary Tumor PC: Pancreatic Cancer Cell Line

Gene Name	Human mRNA for clongation factor 1-alpha	al protein S12.	ıtin 18.	Homo sapiens metallopanstimulin (MPS1)	NA for mucin.	H.sapiens FRGAMMA mRNA (819bp) for folate receptor	H.sapiens mRNA for lung amiloride sensitive Na+ ch	Human FR-gamma' mRNA, complete cds.	Human folate receptor 3 mRNA, complete cds.	somal protein	ye02f02.rl Homo sapiens cDNA clone 116571 5'.	H.sapiens ribosomal protein L37a.	al protein S16	n beta 10	H.sapiens mRNA for ribosomal protein L31.	al protein L27a	mal protein L11.	al protein S6	Human ribosomal protein S28 mRNA, complete cds	Human mRNA for ribosomal protein L17	al protein L35	Human acidic ribosomal phosphoprotein P0	Human M2-type pyruvate kinase mRNA, complete cds.	Human TCB gene encoding cytosolic thyroid hormone-	1,000
	Human mRNA	Human ribosomal protein S12	Human cytokeratin 18.	Homo sapiens m	H.sapiens B1 mRNA for mucin	H.sapiens FRGA	H.sapiens mRN/	Human FR-gam	Human folate re	Human L41 ribosomal protein	ye02f02,r1 Hom	H.sapiens riboso	Human ribosomal protein S16	Human thymosin beta 10	H.sapiens mRN/	Human ribosomal protein L27a	H.sapiens ribosomal protein L1	Human ribosomal protein S6	Human ribosom	Human mRNA	Human ribosomal protein L35	Human acidic ril	Human M2-type	Human TCB ger	11
Accession	69891X	XS3505	X12883	L19739	X83412	Z32564	X76180	U08470	U08471	S64030	T91925	66999X	M60854	M92381	18169X	U14968	X79234	103537	U58682	X52839	U12465		M23725	M26252	111117
PC	412	125	502	179	102					358	0	252	801	203 231	191	142	49	120	2	9	184	145	55		33.
P	136	265 105	36	74	99					179 104 358	48	55	9	203	38	65	23	28	44	4	22	27	46		٠.
NC TU CL PT PC	487 136 412		245	107	186					(179	9/1	172	138	136	133	711	Ξ	109	103	<u></u>	8	68	8		6
TU	62	99	83	53	48					128	0	19	39	90	37	ES	64	39	59	48	43	15	30		٤
S	71	72	137	S	31					115	0	59	20	8	30	38	42	56	32	36	39	22	70		30
Tag Number	H978825	H615043	H263478	H278636	IH					H1027448	H906438	H33979	H374027	H696375	H41531	H567488	H424694	H618199	H549145	H857362	H416106	H475448	H955718		11160100
Tag Segmence	CATGTGTGTTGAGAG	CATGGCCGAGGAAGG	CATGCAAACCATCCA	CATGCACAAACGGTA	CATGAAAAAAAAA					CATGITIGGICCICIG	CATGTCTCCATACCC	CATGAAGACAGIGGC	CATGCCGTCCAAGGG	CATGGGGGAAATCGC	CATGAAGGAGATGGG	CATGGAGGGAGTTTC	CATGCGCTGGTTCCA	CATGGCCGTGTCCGC	CATGGACGACACGAG	16 CATGTCACCCACACC	17 CATGCGCCGCCGGCT	CATGCTCAACATCTC	CATGTGGCCCCACCC		
-		, ,		1	1	ì	1	\top	T	9	1	~		1_		2	2	=	13	٥	Ļ	<u>∞</u>		1	7

-	21 ICATGAGCATCTCCAG	H150997	0	0	77	0	0	H09058	yl96f11.r1 Homo sapiens cDNA clone 45943 5'.
.1								Z44640	H. sapiens partial cDNA sequence; clone c-26b05.
1								N75111	y229e01.rl Homo sapiens cDNA clone 284472 5.
1.	CATGGCCTGTATGAG	H621369	24	32	11	33	86	M31520	Human ribosomal protein S24 mRNA.
2	CATGAGCTCTCCCTG	H161624	33	39	9/	21	29	X53777	Human L23 mRNA for putative ribosomal protein.
. 1						<u> </u>	-		gb/AA223340/AA223340 Homo sapiens cDNA clone 650651 3' similar to
24	CATGCCAGGAGGAAT	H338081	27	12	77		-	AA223340	gb:Y00371 mai HEAT SHOCK COGNATE 71 KD PROTEIN (HUMAN)
25	CATGGCCAAGCCCCA	H672342	30	55	2		2	U12404	Human Csa-19
	CATGAGGAAAGCTGC	H163999	31	42	2	32	146	F16378	H.sapiens EST sequence (135-18) from skeletal muscle
	CATGAACGCGGCCAA	H26261	29	46	69	24	62	Z23063	Homo sapiens macrophage migration inhibitory factor
	CATGCCAGAACAGAC	H335945	23	39	99	42	148	X79238	H.sapiens ribosomal protein L30.
	CATGGCCGCCATCTC	H615736	7	01	9	\vdash	22	US5017	Human transketolase (TKT)
2	CATGGTGTTAACCAG	H769045	16	16	65	17	92	L25899	Human ribosomal protein L10
=	CATGCCTCGGAAAAT	H383489	6	13	64	23	46	Z26876	H.sapiens ribosomal protein L38.
32	CATGAGGTCCTAGCC	H177610	15	27	63	43	4	X06547	Human class Pi glutathione S-transferase
33	CATGGTTCCCTGGCC	H775658	31	26	63	32	98	X65923	H.sapiens fau mRNA.
×	CATGTAAGGAGCTGA	H796831	32	28	29	42	89	X77770	H.sapiens RPS26
12	CATGAACTAAAAAA	H28673	7	14	09	17	39	W52460	2c45e11.rl Soares senescent fibroblasts NbHSF Homo
1							Н	N92893	zb71h03.s1 Homo sapiens cDNA clone 309077 3'.
36	CATGATTTGTCCCAG	H260949	11	13	57	6	91	X14957	Human hmgl mRNA for high mobility group protein l.
37	CATGATAATTCTTTG	H200576	13	27	53	8	8	U14973	Human ribosomal protein \$29
38	CATGCCCCAGCCAGT	H348756	18	23	53	2	35	UI4990	Human XPIPO ribosomal protein S3 (rpS3)
39	CATGGGAGTGGACAT	H667269	15	13	49	2	45	L11566	Homo sapiens ribosomal protein L18 (RPL18)
8	CATGTAAAAAAAA	H786433	13	œ	48	<u>0</u>	56	H08238	yi87a01.rl Homo sapiens cDNA clone 44932 5'.
1-4	CATGGTGTTGCACAA	H769605	19	21	48	7.	47	X79239	H.sapiens ribosomal protein S13.
42	CATGGCCAGCCAGC	H608595	9	21	47	=	15	U31657	Human unknown protein mRNA, partial cds.
1						H	\vdash	H41030	yn92a10.rl Homo sapiens cDNA clone 175866 5'.
5	CATGGGCTCCCACTG	H685384	14	24	47	23	15	M16660	Human 90-kDa heat-shock protein
44	CATGTCAACTTCTGG	H853983	0	0	46	7	0	N57419	yw82e04.r1 Homo sapiens cDNA clone 258750 5' simil
45	CATGGATGCTGCCAA	H583573	9	12	46	27	8.	X59357	Human mRNA for Epstein-Barr virus small RNAs (EBER)
1						-		L21756	Homo sapiens acute myeloid leukemia associated protein
1								D17652	Human mRNA for HBp15/L22, complete cds.
46	CATGAATAGGTCCAA	H51925	13	31	\$	-	53	M64716	Human ribosomal protein S25
1	CATGGCTTTTAAGGA	H655115	ø	26	45	22	8	\neg	Homo sapiens ribosomal protein S20 (RPS20)
8	CATGAATGCAGGCAG	H58533	7	12	44	9	7.7	M61831	Human S-adenosylhomocysteine hydrolase (AHCY)

[A TOCOCO A COTOCO	H610939	∞	8	43	0	22	221507	Human elongation factor 1 delta (EF 1 delta)
\$ 5	CATGGGCGGCGTTCG	H678334	9	9	2	∞	∞	M13932	Human ribosomal protein S17 mRNA
3 ~	- 1	H928269	4	26	42	15	42	M10036	Human triosephosphate isomerase
: 2	CATGTGTACCTGTA	H968173	14	24	42	35	49	K00558	human alpha-tubulin
2 2		H672265	∞	7	41	12	87	L19527	Homo sapiens ribosomal protein L27 (RPL27)
2	CATGAACTAACAAA	H28737	9	14	40	14	15	X63237	H.sapiens Uba80 mRNA for ubiquitin.
2	_	H837237	0	0	38	0	6		Unknown
×	_	H803369	7	17	38	14	42	X69391	H.sapiens ribosomal protein L.6.
2	CATGGTTAACGTCC	H770486	8	17	38	12	25	H11182	ym14a02.r1 Homo sapiens cDNA clone 47866 5'
;	7						_	T40302	ya31g04.r5 Homo sapiens cDNA clone 62262 5'
						┢		T89480	yd98a05.r1 Homo sapiens cDNA clone 116240 5'
85	CATGGAGACTCCTGC	H558943	13	12	38	32	10	H01362	yi99c06.r1 Homo sapiens cDNA clone 147370 5'
2	CATGATCCACATCG	H217399	3	01	37	10	14	H94371	yw54e05.r1 Homo sapiens cDNA clone 255064 5'.
	_					-		T49412	ya75b09.r1 Homo sapiens cDNA clone 67481 5'.
							-	T51058	yb55a12.rl Homo sapiens cDNA clone 75070 5'.
9	CATGGAAGCTTTGCA	H534522	=	13	37	14	25	X07270	Human heat shock protein hsp86.
3 3	CATGCTGGCGAGCG	H501287	2	6	36	3	18	M91670	Human ubiquitin carrier protein (E2-EPF)
3	Т	H493633	13	8	36	8	56	X74070	H.sapiens transcription factor BTF 3.
15	_	H24951	7	13	35	22	40	V00599	Human beta-tubulin
3		H602783	6	16	35	2	17		H.sapiens mRNA for elongations factor Tu-mitochondria
	7							L38995	Homo sapiens nuclear-encoded mitochondrial elongatation factor
						-	┢	\$75463	P43=mitochondrial elongation factor homolog [human
3,9	CATGCATCTTCACCA	H319302	12	14	35	6	91	H48893	yq80b12.rl Homo sapiens cDNA clone 202079 5'
3 3		H621035	2	~	32	<u>∞</u>	107	X71973	H.sapiens GPx-4 mRNA for phospholipid hydroperoxidase
129	CATGACAGGCTACGG	H76231	0	5	31	64	0	M95787	Human 22kDa smooth muscle protein (SM22)
89		H528067	5	7.1	31	14	25	H80294	yu59g01.s1 Homo sapiens cDNA clone 230448 3'.
		a						R74294	yi57f06.r1 Homo sapiens cDNA clone 143363 5'.
3	CATGGAAGCCAGCCA	H533798	-	3	30	6	11	L36055	Human 4E-binding protein 1
2	CATGTTACCATATC	H988366	10	28	30	16	98	F17005	H.sapiens EST sequence (011-T1-18) from skeletal muscle
=	1	H1023249	-	2	29	-	7	H10519	yl90g04.rl Homo sapiens cDNA clone 45563 5'.
72	$\overline{}$	H874103	0	9	29	0	0		Unknown
2	7-	H246019	∞	6	29	25	26	X04409	Human coupling protein G(s) alpha-subunit
74	1	H298495	2	7	28	-	24	- 1	Human UbA52 adrenal mRNA for ubiquitin-52 amino acid
75	CATGGTTCGTGCCAA	H777109	6	28	28	-	46		H.sapiens EST sequence (005-X3-16) from skeletal m
76	76 CATGGACGTGTGGGC	H552683	3	4	27	2	91	X52317	Human histone H2A.Z.

	CATOOTA A A A A A A A A	H458753	4	8	27	61	∞	M33680	Human 26-kDa cell surface protein TAPA-1
: ?	CATGGGGTTTTATT	H704500	4	-	27	9	∞_	L28809	Homo sapiens dbpB-like protein
8 2	CATGCCGATCACCGG	H363799	7	٥	27	7	15	M29536	Human translational initiation factor 2 beta subunit
1	CATGCACAGAGA	H594051	9	6	56	7	29	W07137	za92a11.rl Soares fetal lung NbHL19W Homo sapiens
3								D20503	Human HL60 3'directed Mbol cDNA, HUMGS01477, clone
								N91592	Soares fetal lung NbHL19W Homo sapiens cDNA clone 303055 31.
									yv84c07.s1 Homo sapiens cDNA clone 249420 3' similar to contains Alu
								H83884	repetitive element;.
:	CATGICITATORA	H908373	7	=	56	=	2	Z22572	H.sapiens CDEI binding protein mRNA.
	כאומוכובושים							L09209	Homo sapiens amyloid protein homologue mRNA, compl
								L19597	Human binding protein mRNA, partial cds.
								S60099	APPH=amyloid precursor protein homolog [human, pla
8	CATGGTTTCCCCAAG	H783697	_	0	22	6	0	W07587	zb06f02.rl Soares fetal lung NbHL19W Homo sapiens
72	200000000000000000000000000000000000000							N28502	yx36f06ir1 Homo sapiens cDNA clone 263843 5'
					Γ		-	N35630	yx62a03.rl Homo sapiens cDNA clone 266284 5'
18	CATOCCTGTCCAGO	H388426	2	3	25	5	13	Z40265	H. sapiens partial cDNA sequence; clone c-1xe03.
	200000000000000000000000000000000000000							W02723	zc65c03.s1 Soares fetal heart NbHH19W Homo sapiens
							\vdash	N24893	yx99h09.s1 Homo sapiens cDNA clone 269921 3'.
						Γ		N32178	yy25b09,s1 Homo sapiens cDNA clone 272249 3'.
5	CATCITCATCATCIGA	H865503	5	15	25	S	7	H21873	yl34b10.s1 Homo sapiens cDNA clone 160123 3' simil
5								H26394	yl48e12.s1 Homo sapiens cDNA clone 161518 3' simil
								H69857	yr88d02.s1 Homo sapiens cDNA clone 212355 3' simil
								H70714	yu69b11.s1 Homo sapiens cDNA clone 239037 3' simil
ě	CATGCCTGCCTTGT	H358783	n	∞	25	16	=	X55110	Human mRNA for neurite outgrowth-promoting protein
3/2		H617048	-	_	24	0	_	X03168	Human mRNA for S-protein.
3									zo32d09.s1 Stratagene colon (#937204) Homo sapiens cDNA clone 588593
8.7	CATGITGGTCAAAAA	H1023233	7		24.	7	7	AA143561	3' similar to contains LTR7.tl LTR7 repetitive element
اة									2001g11.s1 Stratagene colon (#937204) Homo sapiens cDNA clone 566468
								AA152342	
									2186h I i.s I Stratagene colon (#937204) Homo sapiens cDNA clone 51 1557
								AA115727	3' similar to contains LTR7.t1 LTR7 repetitive element
8	CATGCAAAATCAGGA	H262987	9	7	24	S	15	R76502	yi61f09.rl Homo sapiens cDNA clone 143753 5.
3				_				T32681	EST52915 Homo sapiens cDNA 5' end similar to None.
								T34662	EST72468 Homo sapiens cDNA 5' end similar to None.
2	CATGGAAGATGTGGG	H533435	-	S	23	4	7	H04634	lyj49h03.r1 Homo sapiens cDNA clone 152117 5.
;								İ	

			_			-		F00364	H. sapiens partial cDNA sequence; clone 76D12; ver
8	on CATGGTGCTCATTCA	H761150	0	∞	23	9	4		yj21c05.s1 Homo sapiens cDNA clone 149384 3'.
								H84813	yv86c02.s1 Homo sapiens cDNA clone 249602 3' simil
								H84956	yv88f07.s1 Homo sapiens cDNA clone 249829 3' simil
5	CATGGCTITACTITG	H654464	4	S	23	6	5	L38961	Homo sapiens putative transmembrane protein (B5)
8	CATGTTTCTGAAAA	H1046401	9	13	23	01	01	304026	Human thioredoxin (TXN) mRNA
: 5	CATGTTGCTCACACA	H1023250	-	4	22	0	4	D11078	Human RGH2 gene.
8	CATGGATTTCTCAGC	H589267	0	0	22	0	19	X53279	Human mRNA for placental-like alkaline phosphatase
98	95 CATGAGGAGGGAGGC	H166539	7	3	22	2	4	M77836	Human pyrroline 5-carboxylate reductase mRNA,
8	CATGGCTTAACCTGG	H651359		4	22	2	4	X07674	Human glutamate dehydrogenase
9	CATGCTCTTCGAGAA	H490889	4	80	22	27	19	Y00433	Human mRNA for glutathione peroxidase
•		H132098	-	7	71	٥	9	X67951	H.sapiens mRNA for proliferation-associated gene
g		H346761	3	3	21	2	24	U38846	Human stimulator of TAR RNA binding (SRB)
								D16933	Human HepG2 3' region cDNA, clone hmd4f1 I.
9	CATGCACTTCAAGGG	H294155	0	3	20	47	107	U42376	Human retinoic acid induced RIG-E
=	_	H631331	2	3	20	4	_		Unknown
<u> </u>	CATGTTACCTCCTTC	H989024	4	7	20	3	77	F17524	H.sapiens EST sequence (012-T2-32) from skeletal in
[5]	CATGACTCTGCCAAG	H122449	4	7	70	3	,		Unknown
⊒	IN CATGTCAGATGGCGT	H861095	_	9	61	12	7	W52942	zc03h05.r1 Soares parathyroid tumor NbHPA Homo sap
Š	TITTITI CATGGGCCTTITITI	H679936	-	3	61	S	~	R21316	yg48h11,r1 Homo sapiens cDNA clone 35917 5' simila
\ <u>\\\\\\\\\\\\\\\\\\\\\\\\\\\\\\\\\\\\</u>	106 CATGITGGACGCGCTG	H951912	0	0	19	0	0	X00566	Human lipoprotein apoAl.
=	107 CATGCCTGCTCCCTG	H386904	0	5	61	9	~	M80244	Human E16 mRNA
108	CATGGCCACACCCCA(C)	H607318	2	9	18	8-	15	H27927	yl58c11.s1 Homo sapiens cDNA clone 162452 3' simil
60	CATGATTATTTTCT	H249854	2	3	-81	5	20	X57959	H.sapiens ribosomal protein L7.
2	CATGGAACCCTGGGA	H529899	2	7	8	~	2	AA299898	EST12509 Uterus tumor I Homo sapiens cDNA 5' end
Ξ	CATGGGCTGATGTGG	H686319	3	5	∞		2	U09510	Human glycyl-tRNA synthetase.
=	112 CATGTCAATAAAGAA	H855049	3	2	22	4	4	X76013	H.sapiens QRSHs mRNA for glutaminyl-tRNA synthetas
13	CATGAAAGTGAAGAT	H11785	0	7	17	0	S	W16529	zb10a11.ri Soares fetal lung NbHL19W Homo sapiens
								W35192	zc70b05.rl Soares fetal heart NbHH19W Homo sapiens
_								W52451	zc45d09.rl Soares senescent fibroblasts NbHSF Homo
=	114 CATGCACGCGCTCAA	H288373	0	E	12	0	3	D38251	Human mRNA for RPB5 (XAP4)
=	15 CATGAACTAATACTA	H28872	-	9	11	13	31	D52570	Human fetal brain cDNA 5'-end GEN-081G12.
								D52758	Human fetal brain cDNA 5'-end GEN-087A08.
									Human fetal brain cDNA 5'-end GEN-407H12.
116	116 CATGCTGTACCTGGA	H504187		0	11	12	9	M22490	Human bone morphogenetic protein-2B (BMP-2B)

1.1	17 CATGOOACCCCACGC	H398663	2	9	11	48	-	M12529	Human apolipoprotein E
	CATGTAGAAAATAA	H819213	0	-	9	2	7	X16539	H.sapiens RNA for neuroleukin gene.
						-		M27691	Human transactivator protein (CREB) mRNA, complete
61:1	CATGATCTTGAAAGG	H228867	0	0	16	5	3	M86667	H.sapiens NAP (nucleosome assembly protein)
	CATGCAGCTGGCCAT	H302741	0	-	16	14	0	X53743	H.sapiens mRNA for fibulin-1 C.
2	CATGATCTTGAAAGG	H228867	0	0	16	5	3	Z26328	H. sapiens partial cDNA sequence; clone HEC059
121	121 CATGATCTTGAAAGG	H228867	0	0	16	5	3	Z26328	H. sapiens partial cDNA sequence; clone HEC059
122	CATGGTGGAGGTGCG	H762554	7	2	91	3	5		Human 100 kDa coactivator mRNA
123	CATGGTGGACCCCAA	H762197	1	5	15	7	10	R91724	yp98e02.r1 Homo sapiens cDNA clone 195482 5' simil
						_		WS1770	zc48a02.r1 Soares senescent fibroblasts NbHSF Homo
					-	_		N42086	yy05b03.rl Homo sapiens cDNA clone 270317 5'
124	CATGGAGCAGCTGGA	H561787	0	5	15	2	4		yi94c02.r1 Homo sapiens cDNA clone 146882 5'
								R95056	yq44f01.r1 Homo sapiens cDNA clone 198649 5' simil
1.25	CATGGGGGGGCT	H633002 ·	-	9	5	∞	7	F16507	H.sapiens EST sequence (147-09) from skeletal musc
				-		-	_	T50201	yb77h05.r1 Homo sapiens cDNA clone 77241 S' simila
136	CATGATTGGCTTAAA	H256497	-	∞	5	0	16	\$85655	Human prohibitin
122	127 CATGGAAAATITAA	H524541	0	60	5	4	0	M38188	Human unknown protein from clone pHGR74 mRNA, comp
200		H577840	0	2	15	0	0	Y00711	Human lactate dehydrogenase B (LDH-B).
200	139 CATGAGCCTTTGTTG	H155632	-	7	15	23	5	D83174	Human collagen binding protein 2.
120	10 CATGTCTGCACCTCC	H910430	0	0	15	0	2	X70940	H.sapiens elongation factor 1 alpha-2.
2 2	CATGAACAGAAGCAA	H18469	0	7	13	3		T30623	EST19638 Homo sapiens cDNA 5' end similar to None.
					\vdash	-	-		HUMGS0004747, Human Gene Signature, 3'-directed cDNA
								C01011	sequence.
						-	_		zm62d06.s1 Stratagene fibroblast (#937212) Homo sapiens cDNA clone
							▼	AA111865	530219 3
								WS6516	zd16c08,r1 Soares fetal heart NbHH19W Homo sapiens
132	CATGTGTTCAGGACC	H980130	-	-	4	5	11	H30299	yo77d04.r1 Homo sapiens cDNA clone 183943 5' simil
_								H50265	yo28c02.rl Homo sapiens cDNA clone 179234 5.
133	CATGTAGATAATGGC	H822331	-	4	14	9	14	W01702	za37a06.rl Soares fetal liver spleen INFLS Homo sa
								W04495	za58b10.rl Soares fetal liver spleen INFLS Homo sa
								W23528	zc71g11.s1 Soares fetal heart NbHH19W Homo sapiens
134	CATGCTTAATCCTGA	H508767	0	9	14	9	12		Human HepG2 3'-directed Mbol cDNA, clone hm02e09.
135	CATGGGCAGAGGACC	H673954	0	9	14	2	=	ı	H.sapiens nm23H1 gene.
136	CATGTGACTGAAGCC	H925194	0	5	14	3	0		EST85850 Homo sapiens cDNA 5' end similar to None.
							H	T35536	EST86951 Homo sapiens cDNA 5' end similar to None.

T35545 EST87066 Homo sapiens cDNA 5' end similar to None.	H01694 yj33g11.s1 Homo sapiens cDNA clone 150596 3'.	N78851 zb17d08.s1 Homo sapiens cDNA clone 302319 3'.	N78931 za92h06.s1 Homo sapiens cDNA clone 300059 3'.	H90469 yv01e06.rl Homo sapiens cDNA clone 241474 5' simil	R76765 yi63g01.rl Homo sapiens cDNA clone 143952 5' simil	T35045 EST79335 Homo sapiens cDNA similar to None	H51447 yo31a05.r1 Homo sapiens cDNA clone 179504 5'.	W46469 zc32c05.rl Soares senescent fibroblasts NbHSF Homo	W51800 zc48e04.rl Soares senescent fibroblasts NbHSF Homo	R33196 yh77f08.rl Homo sapiens cDNA clone 135783 5.	J04799 Human prothymosin-alpha		U02389 Human hLON ATP-dependent protease mRNA	729819 EST96617 Homo sapiens cDNA 5' end similar to ATP-d	X14850 Human histone H2A.X.	J04088 Human DNA topoisomerase II (top2) mRNA	K01891 Human beta globin retrovirus-like repetitive element	1188396 EST28e05 Homo sapiens cDNA clone 28e05		D28480 Human mRNA for hMCM2, complete cds.	D55716 Human B lymphoma mRNA for P1cdc47, complete cds.	T30327 EST14849 Homo sapiens cDNA 5' end similar to None.	T34394 EST66942 Homo sapiens cDNA 5' end similar to None.	T47475 yb14c03.rl Homo sapiens cDNA clone 71140 5.	T50289 yb14h08.rl Homo sapiens cDNA clone 71199 5'.	Unknown		U33818 Human inducible poly(A)-binding protein	D16891 Human HepG2 3' region cDNA, clone hmd2c11.	M29882 Human epolipoprotein A-II	Z49216 H.sapiens mitoxantrone-resistance associated mRNA.	Unknown	Unknown	M93651 Human set gene
\vdash	-		-	13			6		_	\vdash	01	12	2	_	9	2	0		8		_	=	_	-		_	7	4	12	0	4		0	8
\vdash	2	-		9			7	-	-	-	∞	8	7	-	-	1	0		_		-	9		-		2	2	4	4	0	5	-	0	7
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	0			-			0	-			F	0	-	-	-	0	0		_		T	0		-	T	0	0	0	0	0	0	0	0	0
	H576495			H765573			H961304				H1003313	H515821	H125315		H526495	H269775	H16303		H496114			H53129				H890535	H697495	H329737	H1048113	H977034	H345789	H63325	H548203	H921067
		1	1																														1	

LIST ICATGTCCTTCTCCAC	H884181	0	5	=	14	∞	X15804	Human alpha-actinin.
158 CATGTATCTGTCTAC	H843485	0	1	11	2	3	T19569	609F Homo sapiens cDNA clone 609 similar to SET protein
CATGACGTTCTCTT	H114144	0	0	=	-	17	Z36249	HHEA 18W H. sapiens partial cDNA sequence; clone HEA 18W;
160 CATGCCCTGAGTCAG	H358581	٥	0	=	0	0	AA207189	zq73e07.r1 Stratagene neuroepithelium (#937231)Homo sapiens cDNA clone 647268 S' similar to TR:E16910 E16910 ENDONUCLEASE.;
161 CATGGAATTCCTCGA	H540023	0	3	=	~		92208NI	za98h04.s1 Homo sapiens cDNA clone 300631 3'.
						-		ze90d01.s1 Soares fetal heart NbHH19W Homo sapiens cDNA clone
							AA025809	366241 3'
					-			zs85h05.s1 Soares NbHTGBC Homo sapiens cDNA clone 704313
							AA279492	3,
162 CATGGACGCCGAACT	H550274	0	_	=	9	0		Unknown
_								zk84f04.s1 Soares pregnant uterus NbHPU Homo sapiens cDNA clone
163 CATGGCGGACTGGGG	H631275	0	0	=		0	7	489535 3's similar to SW:A5_XENLA P28824 A5 PROTEIN PRECURSOR
164 CATGGGAACACACAG	H656453	0	-	=	0	2	R48460	yj67c12.rl Homo sapiens cDNA clone 153814 5'.
								zp01c02.rl Stratagene ovarian cancer (#937219) Homo sapiens cDNA
							AA173819	clone 595106 5'
165 CATGTTGCGGAGCCC	H1022502	0	7	=	2	_	L19183	HUMMAC30X Human MAC30 mRNA, 3' end.
-							H61710	yr24a07.s1 Homo sapiens cDNA clone 206196 3'.
							H77330	yul Ifi2.s1 Homo sapiens cDNA clone 233519 3'.
							N69482	za18d05.s1 Homo sapiens cDNA clone 292905 3'.
166 CATGGCAGACATTGA	H598335	0	7	10	4	6	H41078	yp52c11.s1 Homo sapiens cDNA clone 191060 3' simil
	H294401	0	_	01	2	0	H04630	yj49g03.rl Homo sapiens cDNA clone 152116 5'.
	H719435	0	0	10	24	0		yi66e12.r1 Homo sapiens cDNA clone 144238 5'.
169 CATGTTCCTCGGGC	H1007018	0	-	10	4	13	R32331	yh68g02.s1 Homo sapiens cDNA clone 134930 3' simil
170 CATGCTGCCGAGCT	-497192	0	8	10	_	9		yd77g07.r1 Homo sapiens cDNA clone 114300 5' simil
171 CATGGTGAAAAAA	H753665	0	2	10	3	7	S77357	transcript chill [human, RF1,RF48 stomach cancer c
172 CATGCTGTGCAGCA	H506149	0	9	2	\$	_	M34338	Human spermidine synthase
173 CATGTAGTTTGTGG	-835515	0	-	0.	0	7	U03911	Human mutator gene (hMSH2)
174 CATGATGTAGTAGTG	H242380	0	5	10	6	7	D55671	Human heterogeneous nuclear ribonucleoprotein
175 CATGGACCCACTACC	H545906	0	-	10	3	_		Human lymphocyte activation antigen 4F2 large subunit
176 CATGAAATAGGTTTT	H12992	0	-	10	9	3	D53402	Human fetal brain cDNA 5-end GEN-108D03.
								yb96f02.r1 Homo sapiens cDNA clone 79035 5'.
							D61243	Human fetal brain cDNA 5'-end GEN-171G06.
							N77240	yv44d02.rl Homo sapiens cDNA clone 245571 5'.
177 CATGCCGGGCGTGGT	H371131	0	0	2		2	T35761	EST90898 Homo sapiens cDNA S' end similar to EST c

178 CATGGACTGAGCTTG	H555168	0	8	2		2	T31901	H555168 0 8 10 3 3 T31901 EST40719 Homo sapiens cDNA 5' end similar to None.
179 CATGAAACGCCCAAT	H6481	0	2	2 10 1	-	3	X98264	X98264 [HSMPP41 H.sapiens mRNA for M-phase phosphoprotein, mpp4, 1523bp
180 CATGATGAGGCCGGG	H232027	0	4	4 10 7	7	-		Unknown
181 CATGGCCCACATCCG(A) H610614 0	H610614	0	6	2	9	2	D87433	0 9 10 6 2 D87433 Human mRNA for KIAA0246 gene, partial cds

Table 1 - Transcripts decreased in colon cancer

Transcripts decreased in only colon primary tumors

compared to normal colon (51 genes)

NC: Normal Colon
TU: Colon Primary Tumor
CL: Colon Cancer Cell Line
PT: Pancreatic Primary Tumor
PC: Pancreatic Cancer Cell Line

Is suburned and su	S Isubun Seen DNA RIVa Ite c Island	cession Gene Name 51 Human mRNA for beta-actin. 98 Human mRNA for cytoskeletal gamma-actin. 83 Human mRNA for cytoskeratin 18	PC Accession 111 X00351 75 X04098 502 X12883	PT PC Accession 203 111 X00351 80 75 X04098 36 507 X12883	CL PI PC Accession 185 203 111 X00351 130 80 75 X04098 245 36 502 X12883	CT CL P1 PC Accession 110 185 203 111 X00351 61 130 80 75 X04098 83 245 36 502 X12883	C1 CL F1 FC Accession 110 185 203 111 X00351 61 130 80 75 X04098 83 245 36 502 X12883	NC CT CL PT PC Accession 184 110 185 203 111 X00351 170 61 130 80 75 X04098 137 83 245 36 502 X12883	Tag_Number NC CT CL PT PC Accession H654591 184 110 185 203 111 X00351 H468434 170 61 130 80 75 X04098 H763478 137 83 245 34 502 X12883
n lipocortin II mRNA. In mRNA for calcium dependent protease (small so man RNA for calcium dependent protease (small so feat brain cDNA 5'-end GEN-141D02. OZ.11 Soares fetal heart NbHH19W Homo sapien of fetal brain cDNA 5'-end GEN-141D02. What is the construction of the control of the control of the control of the control of	Human Intervation of Special Interval of Special Open Special Open Special Open Special Open Special Open Special Interval of Special Interval Interval of Special Interval Int	sion	PC Accession 111 X00351 75 X04098	203 111 X00351 80 75 X04098	CL P1 PC Accession 185 203 111 X00351 130 80 75 X04098 248 256 257 X13022 248 256 257 X13022 258	CT CL PI PC Accession 110 185 203 111 X00351 61 130 80 75 X04098 92 748 26 607 V13803	C1 CL F1 FC Accession 110 185 203 111 X00351 61 130 80 75 X04098 92 748 26 503 V13803	NC CT CL PT PC Accession 184 110 185 203 111 X00351 1170 61 130 80 75 X04098 137 823 X13082	Tag_Number NC CT CL PT PC Accession H654591 184 110 185 203 111 X00351 H468434 170 61 130 80 75 X04098 11552479 137 62 246 25 503 V13623
n mRNA for calcium dependent protease (small stens CpG island DNA genomic Mse1 fragment, cl 02.11 Soares fetal heart NbHH19W Homo sapien n fetal brain cDNA 5'-end GEN-141D02. wan thyroid hormone binding protein (p55) mRNA, 05.81 Homo sapiens cDNA clone 270345 3' 05.81 Homo sapiens cDNA clone 270345 3' 05.81 Homo sapiens cDNA clone acyl-CoA deliydrog n mRNA for argininosuccinate synthetase. n mRNA for very-long-chain acyl-CoA deliydrog n keratinocyte cDNA, clone 173. 1 alpha-tubulin mRNA, 3' end. 1 alpha-tubulin mRNA, 3' end. 1 633 EST47188 Fetal kidney II Homo sapiens cCens id1 mRNA. 1 cytochrome c oxidase subunit VIII (COX8) mR 1 n cytochrome c oxidase subunit mRNA, complete Na, K-ATPase alpha-1 subunit mRNA, complete Na, SollR50350 yj59c04.s1 Homo sapiens cDNA clc)	Human mRNA for calcium dependent protease (small subunit) H. Sapiens CpG island DNA genomic Mse1 fragment, cl 2d30d02.1 Soares fetal heart NbHH19W Homo sapiens Human fetal brain cDNA 5'-end GEN-141D02. Unknown Human thyroid homone binding protein (p55) mRNA, yyd5d05.51 Homo sapiens cDNA clone 270345.3' zbb6a05.1 Soares fetal lung NbHL19W Homo sapiens Human mRNA for argininosuccinate synthetase. Human mRNA for very-long-chain acyl-CoA dehydrogen Human keratinocyte cDNA, clone 173. human apha-tubulin mRNA, clone 173. human apha-tubulin mRNA, 3' end. H. sapiens fd1 mRNA. H. sapiens fd1 mRNA. H. sapiens subulit mRNA for subunit mRNA, complete c gblR50350 R50350 yj59c04.s1 Homo sapiens cDNA clone 153		502 X12883 104 D00017	36 502 X12883 53 104 D00017	245 36 502 X12883 36 53 104 D00017	83 245 36 502 X12883 23 36 53 104 D00017	245 36 502 X12883 36 53 104 D00017	83 245 36 502 X12883 23 36 53 104 D00017	H263478 137 83 245 36 502 X12883 H513181 64 23 36 53 104 D00017
iens CpG island DNA genomic Mse1 fragment, cl 02.11 Soares fetal heart NbHH19W Homo sapiens n fetal brain cDNA 5'-end GEN-141D02. wan n thyroid hormone binding protein (p55) mRNA, 05.51 Homo sapiens cDNA clone 270345 3' 05.51 Homo sapiens cDNA clone 270345 3' 05.51 Homo sapiens cDNA clone 270346 3' 05.51 Homo sapiens cDNA clone 173. n mRNA for arginitosuccinate synthetase. n mRNA for very-long-chain acyl-CoA dehydrogen n keratinocyte cDNA, clone 173. 1 alpha-ubulin mRNA, 3' end. 1633 EST47188 Fetal kidney II Homo sapiens cDNA ens id1 mRNA. ens id1 mRNA. ens id1 mRNA. ens id1 subunit mRNA, complete c 0350[R50350 yj59c04.s1 Homo sapiens cDNA clone	H.sapiens CpG island DNA genomic Mse I fragment, cl 2d30d02.rl Soares fetal heart NbHH19W Homo sapiens Human fetal brain cDNA 5'-end GEN-141D02. Unknown Human thyroid hormone binding protein (p55) mRNA, yy65d05.sl Homo sapiens cDNA clone 270345 3' zb06a05.rl Soares fetal lung NbHL19W Homo sapiens Human mRNA for argininosuccinate synthetase. Human mRNA for argininosuccinate synthetase. Human mRNA for very-long-chain acyl-CoA dehydrogen Human alpha-tubulin mRNA, clone 173. human alpha-tubulin mRNA, 3' end. Asapiens idl mRNA, 3' end. H.sapiens idl mRNA for BiP protein. H.sapiens mRNA for BiP protein. H.sapiens mRNA for BiP protein. Human cytochrome c oxidase subunit VIII (COX8) mRNA Human Na,K-ATPase alpha-1 subunit mRNA, complete c gblR50350/R50350 yj59c04.sl Homo sapiens cDNA clone		46 X04106	37 46 X04106	38 37 46 X04106	27 38 37 46 X04106	38 37 46 X04106	27 38 37 46 X04106	H348922 61 27 38 37 46 X04106
02.11 Soares fetal heart NbHH19W Homo sapiens n fetal brain cDNA 5'-end GEN-141D02. ywn 1 thyroid hormone binding protein (p55) mRNA, 05.51 Homo sapiens cDNA clone 270345 3' 55.11 Soares fetal lung NbHL19W Homo sapiens n mRNA for arginiosuccinate synthetase. n mRNA for very-long-chain acyl-CoA dehydrogen n keratinocyte cDNA, clone 173. 1 alpha-tubulin mRNA, 3' end. 1633 EST47188 Fetal kidney II Homo sapiens cDNA ens id1 mRNA. ens id1 mRNA. ens id1 mRNA. 1 cytochrome c oxidase subunit VIII (COX8) mRNa n cytochrome c oxidase subunit mRNA, complete c 1350 R50350 yj59c04.s1 Homo sapiens cDNA clone	zd30d02.11 Soares fetal heart NbHH19W Homo sapiens Human fetal brain cDNA 5'-end GEN-141D02. Unknown Human thyroid hormone binding protein (p55) mRNA, yy65d05.s1 Homo sapiens cDNA clone 270345 3' zb06a05.r1 Soares fetal lung NbHL19W Homo sapiens Human mRNA for argininosuccinate synthetase. Human mRNA for very-long-chain acyl-CoA dehydrogen Human apha-tubulin mRNA, clone 173. human alpha-tubulin mRNA, 3' end. Asa41631 EST47188 Fetal kidney II Homo sapiens cDNA H.sapiens Id1 mRNA. H.sapiens Id1 mRNA. H.sapiens mRNA for BiP protein. Human cytochrome c oxidase subunit VIII (COX8) mRNa Human Na,K-ATPase alpha-1 subunit mRNA, complete c gblR50350lR50350 yj59c94.s1 Homo sapiens cDNA clone		32 Z65513	6 32 265513	42 6 32 265513	4 42 6 32 265513	42 6 32 265513	4 42 6 32 265513	H581974 53 4 42 6 32 Z65513
n fetal brain cDNA 5'-end GEN-141D02. ywn n thyroid hormone binding protein (p55) mRNA, 05.s1 Homo sapiens cDNA clone 270345 3' 55.s1 Homo sapiens cDNA clone 270345 3' 55.s1 Homo sapiens cDNA clone 270345 3' 55.s1 Homo sapiens cDNA, clone 173. n mRNA for argininosuccinate synthetase. n mRNA for very-long-chain acyl-CoA dehydrogen a lapha-tubulin mRNA, 3' end. 1633 EST47188 Fetal kidney II Homo sapiens cDNA ens id1 mRNA. ens id1 mRNA. ens id1 mRNA. n cytochrome c oxidase subunit VIII (COX8) mRNa n cytochrome c oxidase subunit mRNA, complete c 1350 R50350 yj59c04.s1 Homo sapiens cDNA clone 1	n fetal brain cDNA 5'-end GEN-141D02. Dawn n thyroid hormone binding protein (p55) mRNA, 105.s1 Homo sapiens cDNA clone 270345 3' 05.r1 Soares fetal lung NbHL19W Homo sapiens n mRNA for argininosuccinate synthetase. n mRNA for very-long-chain acyl-CoA dehydrogen n keratinocyte cDNA, clone 173. 1 alpha-tubulin mRNA, 3' end. 1633 EST47188 Fetal kidney II Homo sapiens cDNA cens id1 mRNA. ens mRNA for BiP protein. n cytochrome c oxidase subunit VIII (COX8) mRNa n cytochrome c oxidase subunit mRNA, complete c 0350[R50350 yj59c04.s1 Homo sapiens cDNA clone 133030 5'.		32 W61077	6 32 W61077	26 6 32 W61077	22 26 6 32 W61077	26 6 32 W61077	22 26 6 32 W61077	H504098 50 22 26 6 32 W61077
n thyroid hormone binding protein (p55) mRNA, 05.51 Homo sapiens cDNA clone 270345 3' 35.11 Soares fetal lung NbHL19W Homo sapiens n mRNA for argininosuccinate synthetase. a mRNA for very-long-chain acyl-CoA dehydrogen a keratinocyte cDNA, clone 173. a lapha-ubulin mRNA, 3' end. 1633 EST47188 Fetal kidney II Homo sapiens cDNA ens id1 mRNA. ens id1 mRNA. n cytochrome c oxidase subunit VIII (COX8) mRNA ocytochrome c oxidase subunit mRNA, complete c 1846/850350 yj59c04.s1 Homo sapiens cDNA clone 1850/RS0350 yj59c04.s1	n thyroid hormone binding protein (p55) mRNA, in thyroid hormone binding protein (p55) mRNA, in 15.51 Homo sapiens cDNA clone 270345 3' O5.11 Soares fetal lung NbHL19W Homo sapiens in mRNA for argininosuccinate synthetase. In mRNA for very-long-chain acyl-CoA dehydrogen in mRNA for very-long-chain acyl-CoA dehydrogen in mRNA for very-long-chain acyl-CoA dehydrogen in alpha-tubulin mRNA, 3' end. 1633 EST47188 Fetal kidney II Homo sapiens cDNA clons id1 mRNA. ens id1 mRNA. ens id1 mRNA. ens dehydrogen coxidase subunit VIII (COX8) mRNa in Na, K-ATPase alpha-1 subunit mRNA, complete compositions cDNA clone 15301R50350 yj59c04.s1 Homo sapiens cDNA clone 14.r1 Homo sapiens cDNA clone 153030 5'.		4 D60944	18 4 D60944	26 18 4 D60944	15 26 18 4 D60944	26 18 4 D60944	15 26 18 4 D60944	H427848 47 15 26 18 4 D60944
Human thyroid hormone binding protein (p55) mRNA, yyd5d05.s1 Homo sapiens cDNA clone 270345 3' zb06a05.r1 Soares fetal lung NbHL19W Homo sapiens Human mRNA for argininosuccinate synthetase. Human mRNA for very-long-chain acyl-CoA dehydrogen Human keratinocyte cDNA, clone 173. human alpha-tubulin mRNA, 3' end. AA341633 EST47188 Fetal kidney II Homo sapiens cDNA 5' end H.sapiens id1 mRNA. H.sapiens id1 mRNA. H.sapiens alpha-lubulin mRNA, complete c bill man cytochrome c oxidase subunit VIII (COX8) mRNa Human Na,K-ATPase alpha-1 subunit mRNA, complete c gb R50350 R50350 yj59c04.s1 Homo sapiens cDNA clone 153030 3'	n thyroid hormone binding protein (p55) mRNA, 105.s1 Homo sapiens cDNA clone 270345 3' 05.r1 Soares fetal lung NbHL19W Homo sapiens n mRNA for argininosuccinate synthetase. n mRNA for very-long-chain acyl-CoA dehydrogen n keratinocyte cDNA, clone 173. 1 alpha-tubulin mRNA, 3' end. 1633 EST47188 Fetal kidney II Homo sapiens cDNA clons id1 mRNA. 1634 EST47188 Fetal kidney II Homo sapiens cDNA clons id1 mRNA. 1635 EST47188 Fetal kidney II Homo sapiens cDNA clons id1 mRNA. 1639 EST47188 Fetal kidney II Homo sapiens cDNA clone 1350 RS0350 yj59c04.s1 Homo sapiens cDNA clone 14.r1 Homo sapiens cDNA clone 14.r1 Homo sapiens cDNA clone 14.r1 Homo sapiens cDNA clone 153030 5'.	Unknown	8	15 8	21 15 8	10 21 15 8	21 15 8	10 21 15 8	H349801 47 10 21 15 8
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n mRNA for very-long-chain acyl-CoA dehydrogen n keratinocyte cDNA, clone 173. 1 alpha-tubulin mRNA, 3' end. 1633 EST47188 Fetal kidney II Homo sapiens cDNA ens Id1 mRNA. ens Id1 mRNA. ens id4 mRNA, for BiP protein. 1 cytochrome c oxidase subunit VIII (COX8) mRNa ocytochrome c oxidase subunit mRNA, complete c N350 R50350 yj59c04.s1 Homo sapiens cDNA clone 1	n mRNA for very-long-chain acyl-CoA dehydrogen n keratinocyte cDNA, clone 173. 1 alpha-tubulin mRNA, 3' end. 1633 EST47188 Fetal kidney II Homo sapiens cDNA iens Id1 mRNA. ens Id1 mRNA. ens mRNA for BiP protein. n cytochrome c oxidase subunit VIII (COX8) mRNa n cytochrome c oxidase subunit mRNA, complete c 0350[R50350 yj59c04.s1 Homo sapiens cDNA clone 14.r1 Homo sapiens cDNA clone 183030 5'.		10 X01630	2 10 X01630	57 2 10 X01630	6 57 2 10 X01630	57 2 10 X01630	6 57 2 10 X01630	H28235 42 6 57 2 10 X01630
n keratinocyte cDNA, clone 173. 1 alpha-tubulin mRNA, 3' end. 1633 EST47188 Fetal kidney II Homo sapiens cDNA ens Id1 mRNA. ens Id1 mRNA. ens Id1 mRNA for BIP protein. 1 cytochrome c oxidase subunit VIII (COX8) mRNa ocytochrome c oxidase subunit mRNA, complete c N350 R50350 yj59c04.s1 Homo sapiens cDNA clone 1	n keratinocyte cDNA, clone 173. 1 alpha-tubulin mRNA, 3' end. 1633 EST47188 Fetal kidney II Homo sapiens cDNA iens Id1 mRNA. ens Id1 mRNA for BiP protein. ens mRNA for BiP protein. n cytochrome c oxidase subunit VIII (COX8) mRNa n cytochrome c oxidase subunit mRNA, complete c 0350[R50350 yj59c04.s1 Homo sapiens cDNA clone 14.r1 Homo sapiens cDNA clone 14.r1 Homo sapiens cDNA clone 153030 5'.		8 D43682	17 8 D43682	16 17 8 D43682	12 16 17 8 D43682	16 17 8 D43682	12 16 17 8 D43682	H615802 40 12 16 17 8 D43682
1 alpha-tubulin mRNA, 3' end. 1633 EST47188 Fetal kidney II Homo sapiens cDNA ens Id1 mRNA. ens Id1 mRNA. ens Id2 mRNA for BiP protein. n cytochrome c oxidase subunit VIII (COX8) mRNa n cytochrome c oxidase subunit mRNA, complete c N350 R50350 yj59c04.s1 Homo sapiens cDNA clone 1	n alpha-tubulin mRNA, 3' end. 1633 EST47188 Fetal kidney II Homo sapiens cDNA iens Id1 mRNA. ens Id1 mRNA for BiP protein. ens mRNA for BiP protein. n cytochrome c oxidase subunit VIII (COX8) mRNa n cytochrome c oxidase subunit mRNA, complete c 0350[R50350 yj59c04.s1 Homo sapiens cDNA clone 14.r1 Homo sapiens cDNA clone 14.r1 Homo sapiens cDNA clone 183030 5'.		5 D29146	10 5 D29146	36 10 5 D29146	5 36 10 5 D29146	36 10 5 D29146	5 36 10 5 D29146	H960651 40 5 36 10 5 D29146
1633 EST47188 Fetal kidney II Homo sapiens cDNA ens Id1 mRNA. ens Id1 mRNA for BiP protein. n cytochrome c oxidase subunit VIII (COX8) mRNa n cytochrome c oxidase subunit mRNA, complete c N350 R50350 yj59c04.s1 Homo sapiens cDNA clone 1	1633 EST47188 Fetal kidney II Homo sapiens cDNA iens Idl mRNA. ens mRNA for BiP protein. n cytochrome c oxidase subunit VIII (COX8) mRNa n Na,K-ATPase alpha-1 subunit mRNA, complete c 0350 R50350 yj59c04.s1 Homo sapiens cDNA clone 14.r1 Homo sapiens cDNA clone 14.r1 Homo sapiens cDNA clone		39 K00557	6 39 K00557	20 6 39 K00557	10 20 6 39 K00557	20 6 39 K00557	10 20 6 39 K00557	H648575 38 10 20 6 39 K00557
ens IdI mRNA. ens mRNA for BiP protein. n cytochrome c oxidase subunit VIII (COX8) mRNa n Na,K-ATPase alpha-I subunit mRNA, complete c N350 R50350 yj59c04.s1 Homo sapiens cDNA clone I	iens Idl mRNA. ens mRNA for BiP protein. n cytochrome c oxidase subunit VIII (COX8) mRNa n Na,K-ATPase alpha-1 subunit mRNA, complete c 0350[R50350 yj59c04.s1 Homo sapiens cDNA clone		18 AA341633	19 18 AA341633	15 19 18 AA341633	5 15 19 18 AA341633	15 19 18 AA341633	5 15 19 18 AA341633	H955615 37 5 15 19 18 AA341633
ens mRNA for BiP protein. n cytochrome c oxidase subunit VIII (COX8) mRNa n Na,K-ATPase alpha-I subunit mRNA, complete c 3360 R50350 yj59c04.s1 Homo sapiens cDNA clone I	ens mRNA for BiP protein. n cytochrome c oxidase subunit VIII (COX8) mRNa n Na,K-ATPase alpha-1 subunit mRNA, complete c 0350 R50350 yj59c04.s1 Homo sapiens cDNA clone 4.r1 Homo sapiens cDNA clone 153030 5'.		0 X77956	8 0 X77956	36 8 0 <u>X779</u> 56	4 36 8 0 <u>X779</u> 56	36 8 0 <u>X779</u> 56	4 36 8 0 <u>X779</u> 56	H456167 35 4 36 8 0 X77956
n cytochrome c oxidase subunit VIII (COX8) mRNa n Na,K-ATPase alpha-I subunit mRNA, complete c 3350 R50350 yj59c04.s1 Homo sapiens cDNA clone I	n cytochrome c oxidase subunit VIII (COX8) mRNa n Na,K-ATPase alpha-1 subunit mRNA, complete c 0350[R50350 yj59c04.s1 Homo sapiens cDNA clone 14.r1 Homo sapiens cDNA clone 153030 5'.		10 X87949	13 10 X87949	14 13 10 X87949	9 14 13 10 X87949	14 13 10 X87949	9 14 13 10 X87949	H937452 33 9 14 13 10 X87949
1 Na,K-ATPase alpha-1 subunit mRNA, complete c	n Na,K-ATPase alpha-1 subunit mRNA, complete c 0350 R50350 yj59c04.s1 Homo sapiens cDNA clone 4.r1 Homo sapiens cDNA clone 153030 5'.		31 J04823	6 31 J04823	12 6 31 J04823	7 12 6 31 J04823	12 6 31 J04823	7 12 6 31 J04823	H755160 33 7 12 6 31 J04823
350 R50350 yj59c04.s1 Homo sapiens cDNA clone 1	3350 R50350 yj59c04.s1 Homo sapiens cDNA clone 4.r1 Homo sapiens cDNA clone 153030 5'.		13 U16798	9 13 U16798	18 9 13 U16798	5 18 9 13 U16798	18 9 13 U16798	5 18 9 13 U16798	3G H826831 33 5 18 9 13 U16798
	4.rl Homo sapiens cDNA clone 153030 5'.		27 RS0350	19 27 R50350	26 19 27 R50350	7 26 19 27 R50350	26 19 27 R50350	7 26 19 27 R50350	H760267 29 7 26 19 27 R50350

					r	Γ		EST30445 Homo sapiens cDNA 5' end similar to ubiquinol
23 CATGGGGCGCTGTGG	H694767	28	9	20	9	56	T31329	cytochrome-c reductase, 6.4 kDa.
24 CATGCCTCCAGTAC	H382130	27	9	12	3	61		Unknown
25 CATGCCTGTGACAGC	H388627	27	5	14	∞	7	H63643	yr34d11.r1 Homo sapiens cDNA clone 207189 5' simil
26 CATGTCACAGTGCCT	H856806	24	5	80	17	11	W60924	zd27c08.r1 Soares fetal heart NbHH19W Homo sapiens
22 CATGAATAAAGGCTA	H49320	23	5	7	=	-	L25081	Human GTPase (rhoC) mRNA, complete cds.
28 CATGITGITGIAA	H1031929	23	5	13	15		D45887	Human mRNA for calmodulin, complete cds.
29 CATGAAGGTAGCAGA	H44179	23	4	10	91	12	N62815	yy66b11.s1 Homo sapiens cDNA clone 278493 3'.
30 CATGGTGTTGGGGGT	H769707	21	2	5	4		R68653	yi14b06.s1 Homo sapiens cDNA clone 139187 3'.
31 CATGTGCAGCGCCTG	H936344	21	1	5	7	13	X90858	H.sapiens mRNA for uridine phosphorylase.
32 CATGATGGCACGGAG	H238697	20	2	4	0	3	H19458	yn54c02.s1 Homo sapiens cDNA clone 172226 3' simil
11 CATGGCCAGACACCC	H608326	70		9	-	6	T30468	EST17149 Homo sapiens cDNA 5' end similar to None.
13 CATGCTTCTTGCCCC	H515990	70	0	17	3	0	V00491	Human gene for alpha 1 globin.
35 CATGACCCACGTCAG	H86453	61	7	7	22	6	X51345	Human jun-B mRNA for JUN-B protein.
36 CATGGGCTGCCTGCC	H686458	8	3	4	5	8	R72429	yj90e08.s1 Homo sapiens cDNA clone 156038 3'.
						=	R48449	yj67b10.s1 Homo sapiens cDNA clone 153787 3'.
					-		R52128	yj72b03.s1 Homo sapiens cDNA clone 154253 3'.
17 CATGGAGGGCCGGTG	HS67660	81	2	41	9	16	X12910	Human Na+,K+ ATPase gene exons 1 - 3 (alpha III is
38 CATGGATGAATCCGG	H581847	=	-	3	2	7		Unknown
39 CATGAGGCCGACCAC	H153109	9]	7	=	7	5	X81006	H.sapiens HCG I mRNA.
40 CATGGTTCAGCTGTC	H774780	91	7	12		12	L08666	Homo sapiens porin (por) mRNA, complete cds and tr
LI CATGCCTCGCTCAGT	H383443	91	-	∞	9	1	U04627	Human 78 kDa gastrin-binding protein mRNA, complet
12 CATGCAAATAAAGT	HZ65219	15	-	œ	6	0	U17077	Human BENE mRNA, partial cds.
41 CATGTGCGCCCGCA	H940378	15		8	0	3	U28369	Human semaphorin V mRNA, complete cds.
44 CATGGCAGTGGCCTC	H601752	15	0	9	4	3	D12038	Human HepG2 3'-directed Mbol cDNA, clone \$150.
45 CATGCTGGGCCTGAA	H502137	14	0	3	3	18	U77396	Human TNF-alpha inducible responsive element mRNA,
46 CATGGCCCATTGGAG	H611305	13	_	9	13	17	Z29093	H.sapiens EDDR1 gene for receptor tyrosine kinase.
47 CATGAAGAAACCTC	H32792	12	0	2	2	0	T94990	ye38a04.s1 Homo sapiens cDNA clone 119982 3'.
						-	N69310	za25g05.s1 Homo sapiens cDNA clone 293624 3'.
					-	_		2b86e03.s1 Soares senescent fibroblasts NbHSF Homo sapiens cDNA
							N98502	clone 310492 3'
48 CATGGAATGATTTCT	H538878	12	0	9	9	7	F18838	H.sapiens EST sequence (007-X1-01) from skeletal m
								zr21b10.s1 Stratagene NT2 neuronal precursor 937230 Homo sapiens
49 CATGGCCTGGTCCTT	H621272	12	0	3	3		28	cDNA clone 664027 3'
50 CCATGGCCCACACAG	H610579	-	0	-	_	0	M60047	Human heparin binding protein (HBp17) mRNA
			ĺ			i		

3 2 W52456 zc45e09.rl Soares senescent fibroblasts NbHSF Homo 51 CATGGGATTCCAGTT

Transcripts decreased in both colon primary tumors and colon cancer cell lines compared to normal colon (130 genes)

NC: Normal Colon
TU: Colon Primary Tumor
CL: Colon Cancer Cell Line
PT: Pancreatic Primary Tumor

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Cancer	
 DC. Bancrestic Cancer Cell Line	ָ ֭֭֭֭֭֭֭֭֭֭֭֭֚֭֓֞֝֡֡֝
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Gene Name	X12882 Human mRNA for cytokeratin 8.	F15636 [H.sapiens mitochondrial EST sequence (002715)	Unknown	F16940 [H.sapiens mitochondrial EST sequence (009-T1-21) f	M10050 Human liver fatty acid binding protein (FABP) mRNA	S61953 c-erbB3=receptor tyrosine kinase {alternatively sp	F15506 H.sapiens mitochondrial EST sequence (1-1-02) from	T39321 ya04c01.r2 Homo sapiens cDNA clone 60480 5'.	H24673 yl41a01.s1 Homo sapiens cDNA clone 160776 3;	HUMGS02706 Human colon 3'directed Mbol cDNA, HUMGS02706,	clone cm1673.	T96160 ye09b02.s1 Homo sapiens cDNA clone 117195 3'.	X64364 H.sapiens mRNA for M6 antigen.	M11146 Human ferritin H chain mRNA, complete cds.	L15203 Human secretory protein (P1.B) mRNA, complete cds.	X93036 H.sapiens mRNA for MAT8 protein.	yv07h09.rl Homo sapiens cDNA clone 242081 5' similar to SP:A39484	H93844 A39484 ANDROGEN-WITHDRAWAL APOPTOSIS PROTEIN RVPI,	F17001 [H.sapiens mitochondrial EST sequence (011-T1-13) f	Y00503 Human mRNA for keratin 19.	2b05a11.rl Soares fetal lung NbHL19W Homo sapiens cDNA clone	301148 5' similar to gb: V00567 BETA-2-MICROGLOBULIN	W16632 PRECURSOR (HUMAN);	zo31h04.s1 Stratagene colon (#937204) Homo sapiens cDNA clone	588535 3'
Accession	X12882	F15636			M10050	\$61953	F15506	T39321	H24673		D25586	T96160	X64364	M11146	L15203	X93036		H93844	F17001	Y00503					AA 143804 588535 3'
PC	663	497	1	235	0	13	204	0					178	369	3	61		39	139	219			139		
Ы	136	142	2	43	0	30	71	7					144	235	11	32		40	30	57			340		-
CL	304	402	2	93	4	27	132	0					148	84	0	9		97	101	76			7		
TU	191	282	28	348	92	108	242	131					88	75	120	89		24	66	33			178 110 14 340		
NC	803	208	705	512	504	486	367	276					256	202	201	961		194	190	189			178		
Tag Number NC TU CL PT PC Accession	H382109	H460926	H610997	H90022	H81583	H622680	H153361	H545828					H617195	H1026814	H479577	H600670		H224923	H271574	H544012			H782013		
Tag Sequence	CATGCCTCCAGCTAC	2 CATGCTAAGACTTCA	3 CATGGCCCAGGTCAC	4 CATGACCCTTGGCCA	5 CATGACATTGGGTGA	6 CATGGCGAAACCCTG	CATGAGCCCTACAAA	CATGGACCCAAGATA					CATGGCCGGGTGGGC	10 CATGTTGGGGTTTCC	11 CATGCTCCACCCGAA (or G)	12 CATGGCAGGGCCTCA		13 CATGATCGTGGCGGG	4 CATGCAAGCATCCCC	15 CATGGACATCAAGTC			16 CATGGTTGTGGTTAA		
#	_	2	<u></u>	-7	S	0	1	00					6	2	Ε	2		=	14	2			16	L	

L							T		97 +197hf7 s1 Stratagene colon (#937204) Homo sapiens cDNA clone
								AA133597 512115 3	512115.3
L								T53199	T53199 ya86c05.s1 Homo sapiens cDNA clone 68552 3'.
15	7 CTAGTGCTCCTACCC	H947654	174	22	-	0	0	R00081	R00081 ye73c04.s1 Homo sapiens cDNA clone 123366 3'.
1=	18 CATGCACCCTGATG	H284132	172	33	56	5	9	M16364	M16364 Human creatine kinase-B mRNA, complete cds.
1_									yf22e12.s1 Homo sapiens cDNA clone 127630 3' similar to contains Alu
-	19 CATGCCGCTGCACTC	H368200	163	40	4	2	4	R09410	repetitive element
						\vdash	_		HUMGS0003915, Human Gene Signature, 3'-directed cDNA
	·							C01918	sequence.
							-		yq04h09.s1 Homo sapiens cDNA clone 196001 3' similar to
							-	R92735	contains Alu repetitive element
L						-	-		zh78e12.s1 Soares fetal liver spleen INFLS SI Homo sapiens
								W90374	cDNA clone 418222 3' similar to contains Alu repetitive element
2	20 CATGCTGGCCCTCGG	H501111	163	2	0	56	_	X52003	H.sapiens pS2 protein gene.
7	21 CATGCCCCTGGATC	H350116	091	6	24	-	181	M18981	Human prolactin receptor-associated protein (PRA)
3	22 CATGTTCACTGTGAG	H1001401	160	34	13	74	7.1	M64303	M64303 Human galactoside-binding protein mRNA.
12	23 CATGATTGGAGTGCT	H256186	155	34	_	_	9	X16455	X16455 Human mRNA for carcinoembryonic antigen pCEA80-11.
24	24 CATGCTGACCTGTGT	H493039	149	44	32		37	U14943	U14943 Human MHC antigen (HLA-B) mRNA, complete cds.
12	25 CATGAGCAGATCAGG	H149715	145	20	88	156	130	M81457	Human calpactin 1 light chain mRNA, complete cds.
7	26 CATGGGAAAACAGAA	H655433	126	37	0	24	91	C21047	HUMGS0002546, Human Gene Signature, 3'-directed cDNA sequence
				Г		_	-		zo21h08.s1 Stratagene colon (#937204) Homo sapiens cDNA
							*	1A132779	AA132779 clone 587583 3' similar to SW:LEG4_RAT P38552 GALECTIN-4
						-			zl68h06.s1 Stratagene colon (#937204) Homo sapiens cDNA
							_	1A054072	AA054072 clone 509819 3'
				_	L		-		zo18g08.s1 Stratagene colon (#937204) Homo sapiens cDNA clone
								1A132736	AA 132736 587294 3' similar to SW:LEG4 RAT P38552 GALECTIN-4
27	27 CATGTCACCGGTCAG	H857781	122	7	7	30	7	X04412	X04412 Human mRNA for plasma gelsolin.
78	CATGTGCAGCACGAG	H936217	122	56	32	84	2	X77658	X77658 H. sapiens mRNA for HLA-B*7301.
				-	_	_	_		zo35c09.s1 Stratagene colon (#937204) Homo sapiens cDNA clone
52	CATGGGAACTGTGAA	H657337	115	7		4	7 7	21 AA146606 588880 3'	588880 3'
					-				zo35g09.s1 Stratagene colon (#937204) Homo sapiens cDNA clone
					1		-	AA 146775 588928 3'	588928 3'
								2074g11.s	zo74g11.s1 Stratagene pancreas (#937208) Homo sapiens cDNA clone
				٦	1	1	+	14101045	27.6010.3

L				Г		-			zl83f08.s1 Stratagene colon (#937204) Homo sapiens cDNA clone
							~	AA088704 511239 3'	511239 3'
٦	30 CATGCGAGGGGCCAG	H404117	114	32	54	09	40	H00427	lyj23g11.rl Homo sapiens cDNA clone 149636 5'.
						<u> </u>	 .		2063d03.s1 Stratagene pancreas (#937208) Homo sapiens cDNA clone
							9	AA158715 591557 3"	591557 3'
\perp				_		-		T08562	EST06454 Homo sapiens cDNA clone HIBBG31 3' end.
				-	-	_			zm21a12.s1 Stratagene pancreas (#937208) Homo sapiens cDNA clone
							4	AA078845 526270 3"	526270 3'
=	31 CATGTAAATTGCAAA	H790417	113	9	_	0	0	X73502	X73502 H. Sapiens mRNA for cytokeratin 20.
: 2	1) CATGGGTGGGGCC	H686762	113	36	-	45	43	103191	Human profilin mRNA, complete cds.
1 =	33 CATGGTGCTGAATGG	H761359	109	20	30	1 19	Ξ	U02629	U02629 Human smooth muscle myosin alkali light chain mRNA
7	14 CATGGTGCACTGAGC	H758243	107	13	36	34	82	X07059	X07059 Human M4-50 mRNA for HLA class I antigen.
14	15 CATGITTAACGGCCG	H1032614	107	31	14	3	37	F15592	H.sapiens mitochondrial EST sequence (001 T24) from
1						-			zl74e07.sl Stratagene colon (#937204) Homo sapiens cDNA clone
39	16 CATGCCCTCCCGAAG	H357729	901	1.1	7	3	9	1A053660	AA053660 510372 3' similar to contains Alu repetitive element
									HUMGS04077 Human colon 3'directed Mbol cDNA, HUMGS04077,
		-						D25711	cione cm1210
					-	-	-		H.sapiens CpG DNA, clone 140c4, reverse read cpg 14(Mitochondria
3.7	CATGAGGTGGCAAGA	H178755	105	15	77	4	27	Z56800	EST
, e	38 CATGATACTCCACTC	H204104	102	=	0	0	0	M95174	M95174 Human guanylin mRNA, complete cds.
2	39 CATGCTCGCGCTGGG	H484987	2	25	2	4	16		Unknown
						-			yn01b01.r1 Homo sapiens cDNA clone 167113 5' similar to SP:ZK783.1
40	40 CATGGGGGCAGGGCC	H697514	82	32	28	37	65	R90863	CE00760;
!						_		T24702	EST277 Homo sapiens cDNA clone 10H4.
4	CATGGAAGCAGGACC	H533666	8	33	42	28	87		H.sapiens mRNA for non-muscle type cofilin.
£		H338569	75	72	78	30	91	X67325	H.sapiens p27 mRNA.
2		H70211	74	31	30	01	31	F16604	H.sapiens mitochondrial EST sequence (009T28) from
									za16a03.s1 Homo sapiens cDNA clone 292684 3' similar to contains Alu
44	CATGAGAATAGCTTG	H134304	69	29	-	3	0	N69361	اید
<u></u>									ze30b10.s1 Soares retina N2b4HR Homo sapiens cDNA clone
							~	4A015918	AA015918 360475 3' similar to contains Alu repetitive element
		,							y114h01.s1 Homo sapiens cDNA clone 158257 3' similar to contains Alu
								H26689	repetitive element; contains TARI repetitive element ;.
									zr79h11.s1 Soares NhHMPu S1 Homo sapiens cDNA clone 681957 3'
45	45 CATGCGCTGTGGGGT	H424875	88	6	9	5	7 23	4A256365	23 AA256365 similar to WP:C33A12.7 CE05353

L					Γ	r			zc39e11.s1 Soares senescent fibroblasts NbHSF Homo sapiens cDNA
		The state of the s						W47357	clone 324716 3'
									2b90f03.s1 Soares senescent fibroblasts NbHSF Homo sapiens cDNA
								W19276	clone 310877 3'
1								R07159	R07159 yf13h12.s1 Homo sapiens cDNA clone 126791 3'.
46	CATGCATAGGTTTAG	H314109	89	2	0	0	0	L02785	L02785 Homo sapiens colon mucosa-associated (DRA) mRNA
15	47 CATGGCCGACCAGGT	H614731	65	61	0	3	9	U11862	Human clone HP-DAO1 diamine oxidase
48	CATGAGCTCTTGGAG	H161769	64	=	_	-	-	N93240	zb68b06.s1 Homo sapiens cDNA clone 308723 3'.
1									NIB1986 Normalized infant brain, Bento Soares Homo sapiens cDNA
							•	T16906	3'end.
1									yu22h07.s1 Homo sapiens cDNA clone 234589 3' similar to
							-	H78256	SP:SBP_MOUSE P17563 SELENIUM-BINDING
1						 	_		EST47523 Homo sapiens cDNA 3' end similar to similar to Selenium-
							•	T32362	binding protein, liver.
10	49 CATGCCCAACGCGCT	H344474	57	-	0	3	0	V00493	V00493 Human messenger RNA for alpha globin.
19	50 CATGGACGCGCGCG	H550554	55	21	2	7	14		Unknown
	SI CATGACCCCCCCCC	H87386	54	91	15	15	3 2	X51346	X51346 Human jun-D mRNA for JUN-D protein.
10	52 CATGATGCGGGAGAA	H236169	52	9	01	11	7	R34039	R34039 [yh83f04.r1 Homo sapiens cDNA clone 136351 5.
ŧl						-	_	1966011	yj44e07.s1 Homo sapiens cDNA clone 151620 3.
1				Г			_	R33498	R33498 yh83f04.si Homo sapiens cDNA clone 136351 3'.
1						 	_		zl71e06.rl Stratagene colon (#937204) Homo sapiens cDNA clone
~	53 CATGTCAGCTGCAAC	H862097	15	9	0	0	۷ 0	A053043	AA053043 510082 5'
17	54 CATGGTAAGTGTACT	H723890	50	14	15	-	30	F17394	F17394 [H.sapiens mitochondrial EST sequence (007T13) from
150	SS CATGTGTGGTGCTG	H977640	49	70	1.1	21	8	Z13009	H.sapiens mRNA for E-cadherin.
19	56 CATGGCTGTGCCTGG	H650847	48	17	15	8		X15505	X15505 Human mRNA for pancreatic trypsinogen III.
11	57 CATGTGAGTGACAGA	H929299	48	4	0	0	0	H14641	yl26g02.s1 Homo sapiens cDNA clone 159410 3'.
100	S8 ICATGGGCTGGGCCTG	H686744	47	=	13	32	8	M20469	M20469 Human brain-type clathrin light-chain b mRNA,
1									yy92c07.s1 Homo sapiens cDNA clone 281004 3' similar to contains Alu
0	59 CATGTAATCCCAGCA	H800074	46	5	~	∞	=	N50873	N50873 repetitive element; contains element MER32 repetitive element
10	60 CATGGACCAGTGGCT	HS45514	45	-	٥	0	-	U79725	U79725 Human A33 antigen precursor mRNA, complete cds
15	61 CATGGGCACCGTGCT	H673210	44	으	-	7	-		Unknown
12	62 CATGAAGGACCTTIT	H41344	43	2	4	22	24		ym14f06.rl Homo sapiens cDNA clone 47991 5'.
1								H52178	yt85h08.s1 Homo sapiens cDNA clone 231135 3'.
1					-		•	T40539	T40539 ya05b02.s1 Homo sapiens cDNA clone 60555 3'.
1									

AA303091 EST12940 Uterus tumor I Homo sapiens cDNA 3' end	H599903 43 8 17 24 13 W02429 296163 5'.	N20325	N45127 yz13c12.s1 Homo sapiens cDNA clone 282934 3'.		H972720 43 12 14 25 5 U03106 Human wild-type p53 activated fragment-1 (WAFI) mR	H65878 42 16 7 12 11 W37827 clone 322009 3'		W15332 Homo sapiens cDNA clone 322483 3'	ares par	W32410 clone 321378 3'	N32312 Jyw82c01.s1 Homo sapiens cDNA clone 258720 3'.	H828331 41 6 11 6 9 US1478 Human sodium/potassium-transporting ATPase beta-3	H126619 41 7 1 4 35 Unknown	2p44f11.s1 Stratagene muscle 937209 Homo sapiens cDNA clone	7 7 7	R34696 repetitive element;	yh87e04.s1 Homo sapiens cDNA clone 136734 3' similar to contains Alu	R34696 repetitive element;	zq06e03.s1 Stratagene muscle 937209 Homo sapiens cDNA clone AA 1944971628924 3' similar to contains Alu repetitive element	hbc760 Homo sapiens cDNA clone hbc760 3'end similar to nonspacific	H53508 40 12 0 3 0 T11144 crossreacting antigen.	zl67e01.s1 Stratagene colon (#937204) Homo sapiens cDNA clone	AA058357 509688 3' similar to TR:G189087	CO5803 similar to none	H167606 40 11 4 4 5 AA143765 388306 3	zp45b09.s1 Stratagene HeLa cell s3 937216 Homo sapiens cDNA clone
	63 CATGGCAGCTCCTGT				64 CATGTGTCCTGGTTC	A TO O O A A A O CO O A	COLOR DE COL					66 CATGTAGGATGGGG			68 CATGGTAGCAGGTGT						69 CATGAATCACAAATA		47		70 CATGAGGATGGTCCC	

						-		
71 CATGCCAAAGCTATA	H328308	38	Ξ	9	2	18 M3	5252	M35252 Human CO-029.
72 CATGCGGGAGTCGGG	H434907	38	8	9	0	0 R8	R87448	ym89c10.s1 Homo sapiens cDNA clone 166098 3'.
73 CATGGCCGTGGAGAG	H618121	38	6	2	11	26 X7	X79882	H.sapiens Irp mRNA.
74 CATGCCCCGAAGCC	H349706	37	9	0	0	0		Unknown
75 CATGATTTCAAGATG	H259108	37	_	0	0	0 103	103037	Human carbonic anhydrase II mRNA, complete cds.
76 CATGGCCCAGTGGCT	H611050	37	~	-	7	10		Unknown
77 CATGATGGTGGGGGA	H241323	36	7	9	2.5	2 M9	M92843	H.sapiens zinc finger transcriptional regulator mRNA
78 CATGCCTGCCCCCT	H386390	35	12	7	7	9X S		Human ERK1 mRNA for protein serine/threonine kinase
79 CTAGTGGAAAGTGAA	H950457	34	-	1	12	0 0	V01512	Human cellular oncogene c-fos (complete sequence).
80 CATGGTCATCACCAC	H740629	34	٥	0	0	0 03	4279	U34279 Human uroguanylin mRNA, complete cds.
81 CATGCTTATGGTCCC	H511670	34		0	3	I AA2	87021	AA287021 2557c03.s1 Soares NbHTGBC Homo sapiens cDNA clone 701572 3'
or or the state of	H502136	34		4	=	5 TS	T55226 1	yb47a01.s1 Homo sapiens cDNA clone 74280 3' containing L1 repetitive element
מל כאומכוסממכרוכום			1	\dagger	+	+	+-	of Sello el Homo caniens cDNA cione 26129 3' similar to sh: X07173
						22	R37446	INTER-ALPHA-TRYPSIN INHIBITOR COMPLEX COMPONENT II
	•		-					A A AACTOO IN A CONTRACT STATE OF THE CONTRACT
		í	1	1	,	1		
83 CATGGCCCAGGGCCC	H610982	E	~	٥		2 K		Unknown
84 CATGTTTTACTGAT	111047673	33	7	0	4		R81530	yj02b10.r1 Homo sapiens cDNA clone 147547 5'.
							T32348 I	EST47211 Homo sapiens cDNA 3' end similar to None
							'	zd17g02.s1 Soares fetal heart NbHH19W Homo sapiens cDNA clone
						WS	W57810	340946 3'
				-				247e12.s1 Soares ovary tumor NbHOT Homo sapiens cDNA clone
						AA3	98527	AA398527/725518 3'
85 CATGCCTGCTTGTCG	H387054	32	2	-	9	32 X6	X63187 1	H.sapiens HE4 mRNA for extracellular proteinase inhibitor homologue
86 CATGACCTGGGGAGG	H96931	32	9	4	8	9	-	Unknown
								yg52g07.s1 Homo sapiens cDNA clone 36232 3' similar to gb:M33987
87 CATGCCTTCAAATCA	H390158	31	-	0		0 R46	\neg	CARBONIC ANHYDRASE I
88 CATGTCGGAGCTGTT	H893564	ĸ	-	4	-	위	H98618	yx12a06.s1 Homo sapiens cDNA clone 261490 3'.
							.4	zo97h01.s1 Stratagene ovarian cancer (#937219) Homo sapiens cDNA
				1	+	AA.	7,705	AA171705 clone 594865 3'
					-	£	9212	H99212 [yx15g08.s1 Homo sapiens cDNA clone 261854 3'.

15 CATGGGAGGTGGGGC H666539 20 6 5 12 T0344 Habeline genum in RNA, complex cds.	İ						+	-		1.10.12 1. C Shubbit Usms and along
H1003970 30 6 5 32 22 M75161 H1003970 30 7 3 16 17 T30344 H752297 29 1 3 9 3 T60135 H984414 29 5 0 18 0 R23595 H231029 28 5 5 4 6 A4410947 H231029 28 5 5 4 6 A4410947 H286420 28 5 0 5 4 W68230 H286420 28 5 0 5 4 W68230 H578824 27 1 1 24 17 V00594 H510123 27 1 5 9 6 H43742 H538925 27 1 0 2 0 V00497								_ ₹	A029975	470158 3'
H1003970 30 7 3 16 17 T30344 H152297 29 1 3 9 3 T60135 H984414 29 5 0 18 0 R23595 H231029 28 5 4 6 A4410947 H231029 28 5 4 6 AA410947 H236420 28 5 0 5 4 W68230 H286420 28 5 0 5 4 W68230 H510123 27 1 24 17 V00594 H510123 27 1 5 9 6 H43742 H53824 27 1 0 2 0 V00497	68	CATGGGAGGTGGGGC	H666539	30	9	5	32	-	M75161	H.sapiens granulin mRNA, complete cds.
H152297 29 1 3 9 3 T60135 H984414 29 5 0 18 0 R23595 H231029 28 5 4 6 AA410947 H231029 28 5 4 6 AA410947 H236420 28 5 0 5 4 W68230 H286420 28 5 0 5 4 W68230 H578824 27 1 1 24 17 V00594 H510123 27 1 24 17 V00594 H510123 27 1 0 2 0 V00497	S	CATGTTCCACTAACC	H1003970	8	7	3		_	T30344	gblU53204[HSU53204 Human plectin (PLEC1) mRNA, complete cds.
H98414 29 5 0 18 0 R23595 H231029 28 5 4 6 A4410947 H236420 28 5 0 5 4 W68230 H286420 28 5 0 5 4 W68230 H378824 27 1 1 24 17 V00594 H510123 27 1 5 9 6 H43742 H510123 27 1 5 9 6 H43742	5	CATGGGGGGAT	H752297	29	-	3	6			yc22a06.s1 Homo sapiens cDNA clone 81394 3'.
H984414 29 5 0 18 0 R23595 H231029 28 5 4 6 AA410947 H238420 28 5 4 6 AA410947 H286420 28 5 0 5 4 W68230 H578824 27 1 1 24 17 V00594 H310123 27 1 5 9 6 H43742 H518825 27 4 3 1 0 2 0 V00497										gb U67963 HSU67963 Human Iysophospholipase homolog (HU-K5) mRNA
H984114 29 5 0 18 0 R23595 R69445 H231029 28 5 4 6 AA410947 H286420 28 5 0 5 4 W68230 H578824 27 1 1 24 17 V00594 H510123 27 1 5 9 6 H43742 H538925 27 1 0 2 0 V00497								├	1	yh39a12.r1 Homo sapiens cDNA clone 132094 S' similar to gb:D26129
H231029 28 5 4 6 AA410947 H231029 28 5 5 4 6 AA410947 H286420 28 5 0 5 4 W68230 H286420 28 5 0 5 4 W68230 H286420 28 5 0 5 4 W68230 H510123 27 1 24 17 V00594 H510123 27 1 5 9 6 H43742 H5138925 27 4 3 1 0 2 0 V00497	65		H984414	53	~	0	8	\dashv	7	RIBONUCLEASE PANCREATIC PRECURSOR (HUMAN)
H231029 28 5 4 6 AA410947 H231029 28 5 6 4 H02520 H2520 28 5 0 5 4 W68230 H286420 28 5 0 5 4 W68230 H286420 28 5 0 5 4 W68230 H23824 27 1 1 24 17 V00594 H310123 27 1 2 9 6 H43742 H33825 27 4 3 0 0 V00497										yj83c08.s1 Homo sapiens cDNA clone 155342 3' similar to gb:D26129 RIBONUCLEASE PANCREATIC PRECURSOR (HUMAN);.
H231029 28 5 4 6 AA410947 H231029 28 5 4 6 AA410947 H02520 H286420 28 5 0 5 4 W68230 H286420 28 5 0 5 4 W68230 H286420 28 5 0 5 4 W68230 H28824 27 1 1 24 17 V00594 H310123 27 1 5 9 6 H43742 H318825 27 4 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0							\vdash	-	-	yi84h01.s1 Homo sapiens cDNA clone 145969 3' similar to gb:D26129
H231029 28 5 5 4 6 AA410947 H231029 28 5 5 4 6 AA410947 H02520 H286420 28 5 0 5 4 W68230 H286420 28 5 0 5 4 W68230 H286420 28 5 0 5 4 W68230 H310123 27 1 1 24 17 V00594 H310123 27 1 5 9 6 H43742 H310123 27 1 5 9 6 H43742 H310123 27 1 0 2 0 V00497										RIBONUCLEASE PANCREATIC PRECURSOR (HUMAN);.
H231029 28 5 5 4 6 AA410947 H231029 28 5 5 4 6 AA410947 H286420 28 5 0 5 4 W68230 H286420 28 5 0 5 4 W68230 H578824 27 1 1 24 17 V00594 H510123 27 1 5 9 6 H43742 H538925 27 4 3 1 0 2 0 V00497								.x		yj56c03.s1 Homo sapiens cDNA clone 152740 3' similar to gb:D26129 RIBONUCLEASE PANCREATIC PRECURSOR (HUMAN);.
H231029 28 5 5 4 6 AA410947 H286420 28 5 0 5 4 W68230 H286420 28 5 0 5 4 W68230 H578824 27 1 1 24 17 V00594 H510123 27 1 5 9 6 H43742 H538925 27 1 5 9 6 H43742 H51884 27 1 0 2 0 V00497					T		\dagger	\vdash	1	2v35h12.rl Soares ovary tumor NbHOT Homo sapiens cDNA clone
H231029 28 5 5 4 6 AA410947 H286420 28 5 0 5 4 W68230 H286420 28 5 0 5 4 W68230 H578824 27 1 1 24 17 V00594 H510123 27 1 5 9 6 H43742 H538825 27 4 3 1 0 2 0 V00497										755687 5' similar to TR:G459890 G459890 OVEREXPRESSED IN
H286420 28 5 0 5 4 W68230 S C C C C C C C C C C C C C C C C C C		CATGATGACGCTCAC	H231029	28	~	S	4		4410947	TESTICULAR TUMORS
H286420 28 5 0 5 4 W68230 H288420 28 5 0 5 4 W68230 R89822 H578824 27 1 1 24 17 V00594 H510123 27 1 5 9 6 H43742 H518825 27 1 5 0 V00497							\vdash	_	102520	yj40c11.r1 Homo sapiens cDNA clone 151220 5'.
H286420 28 5 0 5 4 W68230 H288420 28 5 0 5 4 W68230 H578824 27 1 1 24 17 V00594 H510123 27 1 5 9 6 H43742 H538255 27 4 3 1 0 2 0 V00497								_		zo12g08.r1 Stratagene colon (#937204) Homo sapiens cDNA clone
H286420 28 5 0 5 4 W68230 S 4 W68230 S 4 W68230 S 4 W68230 S 4 W68230 S 4 W68230 S 4 W68230 S 4 W68230 S 4 W68230 S 4 W68230 S 4 W68230 S 4 W68230 S 4 W68230 S 4 W68230 S 4 W68230 S 5 4 W										586718 5' similar to TR:G459890 G459890 OVEREXPRESSED IN
H286420 28 5 0 5 4 W68230 R89822 H578824 27 1 1 24 17 V00594 H510123 27 1 5 9 6 H43742 H53825 27 4 3 1 0 2 0 V00497								¥	4130551	TESTICULAR TUMORS.
H286420 28 5 0 5 4 W68230 R89822 H578824 27 1 1 24 17 V00594 H510123 27 1 5 9 6 H43742 H33825 27 4 3 1 0 2 0 V00497						П		H		
H578824 27 1 5 9 6 H43742 H510123 27 1 5 9 6 H43742 H5101884 27 1 0 2 0 V00497	3	OT A OTOTO & COTE & C	UCPY8CH	28	٧	¢	٧.			zd33c10.s1 Soares fetal heart NbHH19W Homo sapiens cDNA clone 342450 3' similar to contains Alu repetitive element
H510123 27 1 5 9 6 H43742 H238925 27 4 3 1 0 2 0 V00497	*	CALGORCCIGICATO					1	╁	_	yp90a02.s1 Homo sapiens cDNA clone 194666 3' similar to contains Alu
H510123 27 1 24 17 V00594 H218925 27 1 5 9 6 H43742 H238925 27 4 3 1 0 2 0 V00497										repetitive element;
H578824 27 I I 24 17 V00594 H510123 27 I S 9 6 H43742 H238925 27 4 3 I 0										zk69e08.s1 Soares pregnant uterus NbHPU Homo sapiens cDNA clone
H518824 27 1 1 24 17 V00594 H510123 27 1 5 9 6 H43742 H238925 27 4 3 1 0 H591884 27 1 0 2 0 V00497								_₹	A053322	488102 3' similar to contains element MER6 repetitive element
H510123 27 1 5 9 6 H43742 H238925 27 4 3 1 0 H591884 27 1 0 2 0 V00497	95		H578824	27	_	-	-		V00594	Human mRNA for metallothionein from cadmium-treated cells
H510123 27 1 5 9 6 H45.42 H238925 27 4 3 1 0 H591884 27 1 0 2 0 V00497				1		•	,			yp21d05.r1 Homo sapiens cDNA clone 188073 5' similar to gb:J05021
H238925 27 4 3 1 0 0 V00497 1 1 0 V00497	96	CATGCTTAGAGGGGT	H510123	17	-	1	7	4	7	EZKIN
H591884 27 1 0 2 0	97	CATGATGGCCCATAC	H238925	23	4	-	-			emblY09616 HSICE H.sapiens mRNA for putative carboxylesterase
	86	CATGGCAAGAAGTG	H591884	27	-	0	7		V00497	Human messenger RNA for beta-globin.

8	99 ICATGTACCTCTGATT	H810468	77	2	7	E	12 X6	X65614	H.sapiens mRNA for calcium-binding protein \$100P.
<u> </u> 2	100 CATGATGATGCCACC	H233106	26	0	2	0	2		
									emb Z69881 HSSERCA3M H.sapiens mRNA for adenosine
0	101 CATGTTCTGTAGCCC	H1014566	25	5	0	4	0	-	triphosphatase, calcium
102	102 CATGCCTGTCTGCCA	H388582	24	-	2	Ţ	3 T99568		ye65c02.r1 Homo sapiens cDNA clone 122594 5'.
							T8.	T87539	yd89f09.s1 Homo sapiens cDNA clone 115433 3'.
						-			gb AA347726 AA347726 EST54132 Fetal heart 11 Homo sapiens cDNA
103	103 CATGTATGATGAGCA	H844682	23	4	0	_	0	· ·	5' end similar to transmembrane secretory component
9	104 CATGCTGGCAAAGGT	H500747	23	0	0	0	0		
100	105 CATGCTTGATTCCCA	H517078	23	4	4	17	7 1.42		Homo sapiens bone-derived growth factor (BPGF-1) m
100	106 CATGCTTGACATACC	H516402	22	0	0	7	2 X68	X68277 1	H.sapiens CL 100 mRNA for protein tyrosine phosphase
									Human N-benzoyl-L-tyrosyl-p-amino-benzoic acid hydrolase
107	107 CATGGCTGGCACATT	H649492	77	5	0	0	0 M8	M82962 a	alpha subunit (PPH alpha) mRNA, complete cds
108	108 CATGTCTGAATTATG	H909556	17	_	1	_	1 Xie	X16354 1	Human mRNA for transmembrane carcinoembryonic antigen (CEA)
									H.sapiens mRNA for Gal-beta(1-3/1-4)GicNAcalpha-2,3-
100	109 CATGGGAAGAGCACT	H657554	71	_		3	3 X74	X74570 s	sialyltransferase
								^	yo45d01.s1 Homo sapiens cDNA clone 180865 3' similar to contains
011	10 CATGGCTCTTCCCCA	H646998	70	7	0	_	0 R87	R87768 F	PTR5 repetitive element
								7	yo36g07.s1 Homo sapiens cDNA clone 180060 3' similar to contains
							R85	R85880 F	PTRS repetitive element
	CATGAAATCTGGCAC	1114245	70	2	0	4	3 L20	L20826 H	Human I-plastin mRNA, complete cds.
=	112 CATGTAATTTGCATT	H802708	61	2	0	-	7 Z50	Z50751 I	HSB4BMR H.sapiens mRNA for B4B
						_	U77	7085 F	U77085 Human epithelial membrane protein (CL-20) mRNA, complete cds
							Y07	1 606/	Y07909 HSPAPR H.sapiens mRNA for Progression Associated Protein
\mathbb{E}	113 CATGGTGGGGGCGCC	H764570	<u>«</u>	_	_	8	2 R48	R48529 y	yj64g10.r1 Homo sapiens cDNA clone 153570 5'.
									EST10a24 Clontech adult human fat cell library HL1108A Homo
-	114 CATGTTATGGTGTGA	H998127	17	0	0	_	0 T27		sapiens cDNA clone 10a24.
1.5	HSICATGGGAGAACAGC	H663571	11	_	7	4	0 T86	T86124 y	yd84b04.s1 Homo sapiens cDNA clone 114895 3'.
					_	-		Z	zo15g05.s1 Stratagene colon (#937204) Homo sapiens cDNA clone
		•					AAI	31008 5	AA131008 587000 3
							R49	R49945 y	yj58g11.s1 Homo sapiens cDNA clone 152996 3'.
							TS7	TS7044 y	ya84h01.s1 Homo sapiens cDNA clone 68401 3'.
911	116 CATGCCAACACCAGC	H328787	11	-	0	0	0		
117	17 CATGAGGTGACTGGG	H178299	17	0	0		0		
8	118 CATGGCCATCCTCCA	H609654	91	0	0		0		gbjR73013jR73013 yj94a09.rl Homo sapiens cDNA clone 156376 5.

Ē	HIPICATGITICTCGTCGC	H1039799	15	-	0	4	4	M69013	4 M69013 Human guanine nucleotide-binding regulatory protein
2	POLCATGTCAGAGCGCTG	H860776	15	-	-	_	0		Опкложп
		*							yv72h06.s1 Soares fetal liver spleen INFLS Homo sapiens
									cDNA clone 248315 3' similar to contains element PTR7 repetitive
2	CATOTTCCGCGTTCC	H1006014	14	1	0	0	2	NS8523 element	element
15	CATGTACGGTGTGG	H814011	14	-	0	0	0		Unknown
12	CATGCTCAGAACTTG	H477216	14	0	-	4	13		Unknown
<u>c.</u>	CATGGGACTAAATGA	H662543	23	_	0	_	0	M29540	M29540 Human carcinoembryonic antigen mRNA (CEA), complete cds.
1									HUMGS04154 Human colon 3'directed Mbol cDNA, HUMGS04154,
125	CATGGCTTGGGGATT	H653988	2	0	0	0	_	D25786	D25786 clone cm0215.
									yc36e02.r1 Homo sapiens cDNA clone 82778 5' similar to gb:L07765
								T73613	LIVER CARBOXYLESTERASE PRECURSOR
126	126 CATGACCCAACTGCC	H86138	12	0	0	0	-		Unknown
15	127 CA TGCTGAACCTCCC	H491894	12	0	0	7	2		gb/T95615/T95615 ye40e03.s1 Homo sapiens cDNA clone 120220 3'.
									zr19611.s1 Stratagene NTZ neuronal precursor 937230 Homo sapiens
128	128 CATGCAAGAGTTTCT	H271102	=	0	0	7	0	AA226797	0 AA226797 cDNA clone 663837 3'
1									zq97h01.s1 Stratagene NTZ neuronal precursor 937230 Homo sapiens
								AA218730	AA218730 cDNA clone 649969 3'
									yp57f10.r1 Homo sapiens cDNA clone 191563 5' similar to gb:M90657
129	129 CATGGTCCGAGTGCA	H743610	=	0	0		~	H38178	H38178 TUMOR-ASSOCIATED ANTIGEN L6 (HUMAN);
12	130 CATGTTTGGTTTCAC	H1043445	=	٥	٥	0	0		Unknown

cell lines compared to normal colon (78 genes) Transcripts decreased in only colon cancer

NC: Normal Colon
TU: Colon Primary Tumor
CL: Colon Cancer Cell Line
PT: Pancreatic Primary Tumor
PC: Pancreatic Cancer Cell Line

CATGACACCTAATTGG H285759 CATGAATTTGAGAAGC H260227 CATGTGATTTCACTT H933704 CATGTTCATACACCT H1002566 CATGCCACTGCACTC H314966 CATGCACTAACACCCT H114966 CATGCACTACTCACC H272 CATGCACTACTCAC H272 CATGCAAACATTCTC H1272 CATGCTCATAAGGAA H478249 CATGTCGAAGCCCCC H885334 CATGTCGAAGCCCCC H885334 CATGTCGAAGCCCCC H885334	603 603 5444 4444 5385 603 5444 603 603 603 603 603 603 603 603	1755 411 566 158 556 158 557 134 223 257 4402 223 4446 171 169 98 169 98 183 94 166 91 167 97	161 1 249 1 64 64 1 17 17 17 17 17 17 17 17 17 17 17 17 17 1	333 173 314 191 132 161	F15516 F12396 L08441	H.sapiens mitochondrial EST sequence (1-t-12) H. sapiens partial cDNA sequence; clone c-39e04.
H260227 H933704 H1002566 H335432 H114966 H291282 H1272 H478249 H885334 H103075	┡═┧╌╏╌╏╌╏╸╏ ╾╂╾╂╾╂╾╂╸╂╸┼╸	╶ ┪╸ ┪╸┧╸┨╸┨╸┪╸┪╸┪╸	┈┋┋┋	173 314 191 132 161		H. sapiens partial cDNA sequence; clone c-39e04.
H933704 H1002566 H335432 H114966 H291282 H1272 H478249 H885334 H103075	╎╴╎╶╎╸╎╸╎╸╏╸╏╸╏╸╏╸╏╸	┟╸┩╸╏ ╺┦╶┦╼╂┈╁╼╀╼╂╾╃	├─┤─┤─┤─┤─ ┤─┤─	314 191 132 161		
H1002566 H335432 H114966 H291282 H1272 H478249 H885334 H103075	╏┈╏┈╏┈╏┈╏┈╏┈╏ ┈╏	┡┋┋	╎┤═╎═╏═╏═╏ ═┼	191 132 161		Human autonomously replicating sequence (ARS) mRNA
H335432 H114966 H291282 H1272 H478249 H885334 H103075	╎╴╎╶┤╸╎╸ ┼╸	 - - - - - - - - - - - - - - - - - - -	Ĭ╼╏═╏═╏ ═╂	132	F15553	H.sapiens mitochondrial EST sequence (001714)
H114966 H291282 H1272 H478249 H885334 H103075	╎╌┼╌┼╌╎╸╎╸	▎▕▗┤▗┤▗┤╸┤	┝ ┦╼┦╼╏╼╏ ╼	191	_	Human cortex mRNA containing an Alu repetitive element
	┝╌┼╾┼╌┼╌	╶┤╶┤╌ ┤╾ ┤				H.sapiens mitochondrial EST sequence (141-20)
	├─┤ ╾ ╏╸		- - -	8	7	Human mitochondrion cytochrome b gene, partial cds
			$\dashv \dashv$	223	٦	H.sapiens mitochondrial EST sequence (101-03)
			-	75		H.sapiens mitochondrial EST sequence (1-t-0/)
		+		23		H.sapiens mitochondrial ES1 sequence (022119)
	-	-	1	47	_	yj47a08.s1 Homo sapiens cDNA clone 151862 5.
TTGGCCAGGCT H1025322	-	4	Ξ	21	П	H.sapiens mRNA for MHC class II transactivator.
TTGGTGAAGGA H1027595	98	106 17	183	107		Human thymosin beta-4 mRNA, complete cds.
A TCACGCCCTC H214616	97	186 17	41	49		Human EST overexpressed in pancreatic cancer (xs31)
GCCTGCACCA H941638	67 4	48 25	75	34		Human mRNA for cysteine proteinase inhibitor precursor
AGACCCACAAC H136465	64 1	121 28	24	15		Human fetal brain cDNA 5'-end GEN-129B05.
CATGAGTTTGTTAGT H196339	90	33 17	13	15		Human mRNA for adenocarcinoma-associated antigen
CATGGGAACAACAG H656389	26 4	41 4	33	3	ì	Homo sapiens CD24 signal transducer mRNA
TGGTGTATGCA H965434	53 2	271 6	30	5		Human fetal brain cDNA 3'-end GEN-002A10.
GAAATACAGTT H527436	49	35 10	100	36		Human cathepsin D mRNA, complete cds.
CATGGTGGCTCACGC H763719	49	37 21	72	15		Human Tax I binding protein mRNA, partial cds.
GTGGTGCACAC H765509	45 2	26 18	23	15	\neg	Human metabotropic glutamate receptor i alpha
71 CATGGGTTGGCTTG H704160	44	56 2	9	-	7	tRNASer(UNC) [human, muscle, MERRF/MELAS overlap s
CATGGTGGCGGGTGC H763567	42	32 15	-	5	T48809	yb05c03.rl Homo sapiens cDNA clone 70276 5' contai
25 CATGTAGACTAGCAA H821029	39	23 1	23	2	M69023	Human globin gene.

					,		,	,			,																				_	
Human fetal brain cDNA 3'-end GEN-007C04.	W15552 zb91h11.s1 Soares parathyroid tumor NbHPA Homo sap	H.sapiens mitochondrial EST sequence (132-20) from skeletal muscle	EST186995 HCC cell line (matastasis to liver in mouse) Il Homo	AA315049 sapiens cDNA 5' end	H. sapiens partial cDNA sequence; clone A6A03; ver	yw53h01.s1 Homo sapiens cDNA clone 255985 3'.	Human MHC class I HLA-A2 gene, complete cds.	yf25f12.s1 Homo sapiens cDNA clone 127919 3'.	yl22c10.s1 Homo sapiens cDNA clone 158994 3'.	EST58371 Homo sapiens cDNA 3' end similar to None	H.sapiens mitochondrial EST sequence (129-09)	2154f10.s1 Soares ovary tumor NbHOT Homo sapiens cDNA clone	726187 3'	Z31611.r1 Soares ovary tumor NbHOT Homo sapiens cDNA clone	2467407 el Soares fetal lima NHH 19W Home saniens CDNA clone	308173 3' similar to PIR-A 39484 A 39484 androsen-withdrawal	apoptosis protein RVP1, prostatic - rat	2b19c06.s1 Homo sapiens cDNA clone 302506 3' similar to	PIR:A39484 A39484 androgen-withdrawal apoptosis protein RVP1,	prostatic - rat;	zk39d06.s1 Soares pregnant uterus NbHPU Homo sapiens cDNA	clone 485195 3' similar to PIR:A39484 A39484 androgen-	AA039323 withdrawal apoptosis protein RVP1	Human partial cDNA sequence with CCA repeat region	Human episialin variant A mRNA, 3' end.	Unknown	seq816 Homo sapiens cDNA clone b4HB3MA-COT8-HAP-Ft	H.sapiens mRNA for LI-cadherin.	Homo sapiens huntingtin (HD) gene, exon 66.	dbj[C00470]C00470 HUMGS0007620, Human Gene Signature, 3'-	Land Court of March and Albar Alone 270174 2	No.5551 1yyozgue.si nomo sapiens ciniva cione 2/61/4 5.
D51017	W15552	F16326		AA315049	F01150	N29971	K02883	R09140	R76005	T33596	F16449		AA292959 726187 3'	33400644	AA292400		N92384			N80203			AA039323	U21468	M34088		T10098	X83228	L27415	0.00470	NESCOL	NOSSSI
13	=	6		7	36	7	12	5			92		2	,	7									2	17	0	4	7	2	۳		
25	29	9		8	11	9 :	91	20			14		_		-									20	45	0	3	0	2	-	-	
13	9	=		=	=	0	3	7			7				-									<i>L</i>	0	1	7	7	1	*	•	
144	372	170		13	18	13	14	32			73		6	•	×									218	10	6	11	6	L	4		
38	37	37		33	33	32	32	32			53		78	,	ę									56	52	24	24	22	21	10	7	
H641789	516783H	169669H		H261569	H294488	H386963	H132598	H489822			H609624		H610922	0/0/2011	H956860									H175872	H387596	H188027	H353760	H2235	H607977	04702711	400/01H	
26 CATGGCTAGGTTTAT	27 CATGGGCTTTAGGGA		CATOOOGIA CATOOOGIA	29 CATGATTTCTAAAA	CATGCACTTGCC	_	T	7-	$\overline{}$		34 CATGGCCATGCCTT		15 CATGGCCCAGCGGCC		36 CATGTGGCGCGTGTC									37 CATGAGGGTGTTTTC	38 CATGCCTGGGAAGTG	1-	7	_	1		43 CATGAGGATGTGG	

								2080f04.51 Stratagene ovarian cancer (#937219) Homo sapiens
							AA165679	AA 165679 cDNA clone 593215 3'
					Ι.			zy40a02.si Soares ovary tumor NbHOT Homo sapiens cDNA clone
44 CATGTATAGTCCTCT	H838494	2	-	-	m	4	AA411012 /300/4 3	20014 3
							AA133595 512126 3'	\$12126 3'
			1					2156b12.s1 Soares ovary tumor NbHOT Homo sapiens cDNA clone
							AA292774 726335 3'	726335 3'
AS CATGGGTCTCTT	H710520	20	7	7	7	2	R53216	yj73h02.r1 Homo sapiens cDNA clone 154419 5' simil
45 CATGATGGCTTGAT	H240121	19	4	0	٣	3	D20113	Human HL60 3'directed Mbol cDNA, HUMGS01086, clone
CATGLIGUE	H496981	19	5	0	1	4		Unknown
-	H1013522	61	4	-	∞	2	U35048	Human TSC-22 protein mRNA, complete cds.
48 CATGAAGAAGCAGGG	L	18	4	7	2	8	R81767	yj05g03.r1 Homo sapiens cDNA clone 147892 5'.
49 CATOAGTAGGTGGC	H183018	18	131	2	11	7	D51021	Human fetal brain cDNA 3'-end GEN-00/100/.
SU CATOACAGAGAGAGAGA	1	82	5	~	0	8	D26146	Human DNA for putative protein kinase.
ST CATGGGAAAGTGGT	\perp	18	23	6	5	-		Human alpha-1-antitrypsin mRNA, complete cds.
_	H32926	17	4	0	2	_	R78188	yi81g01.rl Homo sapiens cDNA clone 143680 5.
_	H70965	12	4	0	0	0	M22406	M22406 Human intestinal mucin mRNA, partial cds, clone SM
24 CATOACACCCATOTIC	H144707	17	18	0	0	0	T24507	EST082 Homo sapiens cDNA clone 3E6
CATGAGATCCCA								za63a11.s1 Homo sapiens cDNA clone 297212 3' similar to
							N79237	PIR:S49589 S49589 cortical granule lectin - African clawed frog;
					T		T31354	EST30893 Homo sapiens cDNA 5' end similar to None
	D422214	16	4	0	6	6	H54696	yq92e02.s1 Homo sapiens cDNA clone 203258 3' simil
	0305071	14	0	0	0	0	M22430	Human RASF-A PLA2 mRNA, complete cds.
_	7207571	2	4	2	000	-	AA374631	AA374631 EST86866 HSC172 cells I Homo sapiens cDNA 5' end
S& CAIGGUIIGUIIG	0/245011	<u> </u>			1			zn93g08.r1 Stratagene lung carcinoma 937218 Homo sapiens
							AA137163	AA137163 cDNA clone 565790 5'
								zk10f05.s1 Soares pregnant uterus NbHPU Homo sapiens cDNA
							AA029320	AA029320 clone 470145 3'
A DITTO TO TO TO A	H948543	15	2	0	-	0	D25681	Human colon 3'directed Mbol cDNA, HUMGS04047, clon
-1								2r72g02.s1 Soares NhHMPu S1 Homo sapiens cDNA clone 668978
							AA253331	31
	,						H05110	y175f07.s1 Homo sapiens cDNA clone 43778 3'.
TTOTOTATOTATO	H341720	15	80	-	-	2		Unknown
CATOCCATOOL	11520012	17	23	c	0	0	AA297150	AA297150 EST112734 Colon I Homo sapiens cDNA 5' end
61 CATGGAACAGCICAC	ביחבדרט		1	,				

62 CATGGGGCTACGTCC	H695406	14	4	0	-	٥	M25629	M25629 Human kallikrein mRNA, complete cds, clone clone p
CATGCCCGGCT	H354776	14	7	-	2	7	H18836	ym45d10.s1 Homo sapiens cDNA clone 51262 3'.
								zk01e10.s1 Soares pregnant uterus NbHPU Homo sapiens cDNA
						******	AA026974	AA026974 clone 469290 3'
								zu12c12.rl Soares testis NHT Homo sapiens cDNA clone 731638 5'
-								similar to gb:M61900 Human prostaglandin D synthase gene,
							AA405031	AA405031 complete cds. (HUMAN);
								gb U66894 HSU66894 Human epithelium-restricted Ets protein ESX
64 CATGAGGTACTACTA	H176584	13	6	0	6	∞	U66894	mRNA,
7								Human epithelial-specific transcription factor ESE-1b (ESE-1)
							U73843	mRNA, complete cds
65 CATGCAAATAAATTA	H265232	13	3	0		0	D25996	Human colon 3'directed Mbol cDNA, HUMGS06772
-	H503809	13	9	0				Unknown
\neg								ze88g07.s1 Soares fetal heart NbHH19W Homo sapiens cDNA clone
67 CATGGTTCAATCCCT	H774358	2	6	0	7		AA071520 366108 3	366108 3'
$\overline{}$	-							za90h10.s1 Soares fetal lung NbHL19W Homo sapiens cDNA clone
							N90742	299875 3.
								zn52h06.s1 Stratagene muscle 937209 Homo sapiens cDNA clone
							AA086292 561851 3	561851 3'
68 CATGAATAAAGCCTT	H49304	12	4	0	0	0	D11499	Human HepG2 3'-directed MboI cDNA, clone a-35.
_	H658173	12	7	0	-	0	T16031	IB2474 Homo sapiens cDNA 3'end.
CATGGGATGGC	H670333	12	-	0	9	-	T74426	yc82e01.rl Homo sapiens cDNA clone 22306 5'.
	H715099	12	2	0	2	2	N73771	za61h02.s1 Homo sapiens cDNA clone 297075 3'.
								zh75f08.s1 Soares fetal liver spleen INFLS S1 Homo sapiens cDNA
							W90388	clone 417927 3'
							F03786	H. sapiens partial cDNA sequence; clone c-29h08.
72 CATGTACTGTACTTC	H817952	12	7	٥	0	0	U14631	Human 11 beta-hydroxysteroid dehydrogenase type II
7								ya31a06.s5 Homo sapiens cDNA clone 62194 3' contains Alu
73 CATGCCCTTGCACTC		11	9	-	~	~	T41121	repetitive element,.
_	H440966	11	4	0	7	9		Unknown
75 CATGGCCCCCAACCA	H611590	11	2	0	0	0		Unknown
76 CATGGCCGGCGCTC	H616862	=	2	0	0	0	Z58486	Unknown
77 CATGGGAGGCGCTCA	1	=	_	0	0	0		Unknown

zd42c12.s1 Soares fetal heart NbHH19W Homo sapiens cDNA clone W68073 343318 3' similar to contains Alu repetitive element; 0 Ξ H874226 78 CATGTCCCCGTTACA

Table 4 - Transcripts increased in pancreas_cancer .

	SAGE Tags elevated only in Pancreatic Tumor	NC. Normal Colon	Tu Colon Tumor	CC Colon Cancer Cell Line	PT Pancreatic Tumor	PC: Pancreatic Cell Line	Ca Ba CO E Cit
•	S	ž	F	ŭ	P	7	

	PC Pancreauc Cen Line										
	Tag Sequence	Тад	Tag Number NC Tu CC PT	2	근	<u>5</u>	7	S S		Accession	Gene Name
1	CATGAAAGCAAACCA		H9222	0		1	3	11	Examples R38305	R38305	yh95b04.s1 Homo sapiens cDNA clone 137455 3'
1		-			_	_		L			2k95b03.s1 Soarcs pregnant uterus NbHPU Homo sapiens cDNA clone
		<u> </u>								AA126719	490541 3'
-		_		L	_	L	L				2k51c03.s1 Soares pregnant uterus NbHPU Homo sapiens cDNA clone
										AA044296	486340 3'
T				L	_	L					zl33c08.s1 Soares pregnant uterus NbHPU Homo sapiens cDNA clone
										AA131586	503726 3'
		-			L	L					zo71h12.s1 Stratagene pancreas (#937208) Homo sapiens cDNA clone
2	CATGAAAGCAGTTTA		H9408		_	5	21	3		Examples AA157983	592391 3'
1				L	_	L	L				2t54e04.s1 Soares ovary tumor NbHOT Homo sapiens cDNA clone 726174
										AA292929	3,
		_			L	L					zo78c07.s1 Stratagene pancreas (#937208) Homo zo78c07.s1 Stratagene
										AA159306	pancreas (#937208) Homo
				L	_	L				R54012	yj70h01.s1 Homo sapiens cDNA clone 154129 3'
1		_				_				T62936	yb99f08.s1 Homo sapiens cDNA clone 79335 3'
-	CATGAAAGCGGGGCT		8686H		0	0	0	13	Examples X52426	X52426	H. sapiens mRNA for cytokeratin 13
1	4 CATGAAATCCTGGGT	_	H13803		0		16	7	Examples X51698	X51698	H.sapiens spasmolytic polypeptide (SP) mRNA.
15	SCATGAAATGGACAAC	_	H14865		0	0	0	13	Examples N70419	N70419	za61d12.s1 Homo sapiens cDNA clone 297047 3'
										AA411599	zv16g01.r1 Soares NhHMPu S1 Homo sapiens cDNA clone 753840 5'
		_									11 10 00 THE 12 TO 10 THE 12 T
					_	_				AA410508	zv16gU1.51 Soares NnHMPu Si Homo sapiens cUNA clone 75384U 3
											zl86g12.s1 Stratagene colon (#937204) Homo sapiens cDNA clone 511558
3	6 CATGAACCAGTTTGT		H21247	_	_	9	8	13	Ì	Examples AA115723	3.
		_			_						zo19e04.s1 Stratagene colon (#937204) Homo sapiens cDNA clone 587358
			١	_						AA132875	31
T				L	L						2044a06.s1 Stratagene endothelial cell 937223 Homo sapiens cDNA clone
	-				•					AA147677	589714 3'

						AA206883	240.1012, statisticale and tachton (#25.1252) regions aspicals controlled (#480713)
TICATGAACTCTTGAAG	H30689	3	13	13 17		Examples R51318	yg72f03.s1 Homo sapiens cDNA clone 38681 3'
						T35270	EST82235 Homo sapiens cDNA 3' end similar to None
		-				AA412071	zt65h12.s1 Soares testis NHT Homo sapiens cDNA clone 727271 3'
44744004440	H31221	9	∞	6 130		Examples N63154	yz37f12.s1 Homo sapiens cDNA clone 285263 3'
S CATGMACT GCT T CON	1_		+	<u> </u>		T87236	yc81h04,s1 Homo sapiens cDNA clone 22603 3'
			\vdash	_		AA150720	2146f04.s1 Soares pregnant uterus NbHPU Homo sapiens cDNA clone 5049
			-			AA045773	z168b12.s1 Stratagene colon (#937204) Homo sapiens
UCATGAACTTGGCCAT	H32405	0	0	8 11		Examples X07819	Human pump-1 mRNA homolog, to metalloproteinase,
		,	-			L22523	Human matrilysin gene, exon 5
CATGAAGATCCCCGC	H36183	5 10	14	12 23	Examples R72650	R72650	rj95e05.s1 Homo sapiens cDNA clone 156512 31
							2458e02 st Scares fetal heart NbHH19W Home saniens cDNA clone
							344858 3' similar to SW:CUTA ECOLI P36654 PERIPLASMIC
						W70287	DIVALENT CATION TOLERANCE PROTEIN CUTA
			-	_			1795e05.s1 Homo sapiens cDNA clone 156512 3' similar to
						059628	SPICYCY ECOLI P36634 C-1 YPB CY IOCHROME BIOGENESIS
			+	1			
						,,,,	ap51a11.s1 Stratagene endothelial cell 937223 Homo sapiens cDNA clone
							624668 3' similar to SW:CUTA_ECOLI P36654 PERIPLASMIC
						AA181976	DIVALENT CATION TOLERANCE PROTEIN CUTA
			<u> </u>				Human phosphotyrosine independent ligand p62 for tihe Lck SH2 domain
CATGAAGGGAGGGTC	H43180	1	- 1		_].	1046751	mKNA, complete cds
CATGAAGTTGCTATT	H48756 1	10 9	<u></u>	31 27	Examples 103077	103077	Human co-beta glucosidase (proactivator) mKiNA
			_			M86181	Human prosaposin (PSAP) gene
			-			D00422	Human sphingolipid activator proteins, mRNA
			-	_		103015	Homo sapiens sphingolipid activator protein 1 mRNA
			-			M60255	Human mutant cerebroside sulfate activator protein
1 CATGAATGAAAAA	H57345	10	2	2 10			
TO TO TO TO TO THE STATE OF THE	H66031 1	17 4	24	2 60		Examples N22375	yw37d01.s1 Homo sapiens cDNA clone 254401 3'
			_				zn20e01.s1 Stratagene neuroepithelium NT2RAMI 937234 Homo sapiens
		_	_			A A D 8 4 6 4 3	John A clone 547992 3'

						AA279290	zs84a06.s1 Soares NbHTGBC Homo sapiens cDNA clone 704146 3'
			-			AA046253	zf12a02.s1 Soares fetal heart NbiHH19W Homo sapiens cDNA clone 376682 31
15 CATGACAACTCAATA	H67396	2 7	7 16	37	Examples Z58016	258016	H.sapiens CpG DNA, clone 26c7,
						AA151668	2029c02.s1 Stratagene colon (#937204) Homo sapiens cDNA clone 588290 3' similar to SW:B13_MOUSE P28662 BRAIN PROTEIN 13
			-			W02958	za07e06.rl Soares melanocyte 2NbHM Homo sapiens cDNA clone 291874
16 Chargach CCTGC	H71151		0	2 14	1	zo70e05.s Examples AA1556464 592256 3'	zo70e05.81 Stratagene pancreas (#937208) Homo sapiens cDNA clone 592256 3'
			-			AA025673	ze90h09.s1 Soares fetal heart NbHH19W Homo sapiens cDNA clone
	+		+			T	za89h12.s1 Homo sapiens cDNA clone 299783 3'
17 CATGACCATTGGATT	H85924	8	5 13	4	Examples X02491		Human interferon-inducible mRNA (cDNA 9-27); membrane
			_			104164	Human interferon-inducible protein 9-27 mRNA
							H.sapiens mRNA for interferon-induced 17kDa membra
INCATGACCCTTTAACA	H90050	1 4	2 13	7	Examples X56841		H.sapiens HLA-E gene.
							H.sapiens mRNA for HLA-E heavy chain (exons 4 - 7)
19 CATGACCGCCGTGGT	H91579	49 22	45 70	64	Examples M21186		Human neutrophil cytochrome b light chain p22A
			-				Human p22-phox (CYBA) gene, exons 3 and 4
20 CATGACCTGTGACCA	H97158	0 3	0 28	17	Examples D00244		Human Pro-urokinase gene,
							Human urokinase gene, 3' end
			-				Human pro-urokinase mRNA, complete cds
			_			X02419	Human uPA gene for urokinase-plasminogen activator
21 CATGACGCCCTGCTC	H103912	0	0 11	2	Examples L08835		Human myotonic dystrophy kinase (DM kinase) gene
			_			M87313	Homo sapiens myotonin protein kinase (DM) mRNA
22 CATGACGTGGTGATG	H113380	2 4	4 5	20	Examples H44451		yo75f06.s1 Homo sapiens cDNA clone 183779 3'
							2042f07.81 Stratagene endothelial cell 937223 Homo sapiens cDNA clone 589573 3' similar to SW-L10K RAT 005310 LEYDIG CELL TIMOR 10
						AA157329	KD PROTEIN
			_				2c32g6.s1 Soares senescent fibroblasts NbHSF Homo sapiens cDNA clone 324058 3's similar to SW:L10K RAT 005310 LEYDIG CELL TUMOR 10
						W46455	KD PROTEIN
		1		1			

11 CATGACTCAGCCCGG	H119383	0	0	3 21		Examples M92357		Homo sapiens B94 protein mRNA, complete cds.
	11132631	-	-	5	,,	ZV64875	X64875	Heanjene mRNA for inculin. 11% orough forfor hinding profein 3
24 CATGACTGAGGAAAG	H123211			_L	1	Evanipies	CLOTON	The second to the second descendent incide file second forms hinding
							M31159	rimian growth normane-dependent mount-take growth factor officing protein 3
			-	L			M35878	Human insulin-like growth factor-binding protein-3
			\vdash				S56205	insulin-like growth factor binding protein 3 {3' region}
25 CATGACTGCCCGCTG	H124264	-	0	0 22	0	Examples U65932	U65932	Human extracellular matrix protein 1 (ECM1) mRNA
		\vdash	_	L			U65937	Human extracellular matrix protein 1 (ECM1) gene, exon 9
			-					2003f09.s1 Stratagene colon (#937204) Homo sapiens cDNA clone 566633
26 CATGACTGTATTTC	H126208		4	2	22	Examples	Examples AA148916	3,
2010 110 07		+	-					zo12a11.s1 Stratagene colon (#937204) Homo sapiens cDNA clone 586652
							AA129137	3,
		-	-	L				zi85g09.s1 Stratagene colon (#937204) Homo sapiens cDNA clone 511456
							AA115437	3.
		-	-					zl87e07.s1 Stratagene colon (#937204) Homo sapiens cDNA clone 511620
							AA126967	3.
71 CATGAGCACTGCAGC	H149395	-	2 6	30	16	Examples R24613		yh36c03.rl Homo sapiens cDNA clone 131812
28 CATGAGGAGGGT	H150055	-	0	0	53	Examples H43243		yp05e05.r1 Homo sapiens cDNA clone 186560 5'
29 CATGAGCTGTATTCT	H162622	0	2 0	-	=	Examples X54942		H.sapiens ckshs2 mRNA for Cks1 protein homologue
		-	_					zk50g07.s1 Soares pregnant uterus NbHPU Homo sapiens cDNA clone
10 CATGAGGATGACCCC	H167446	~	7 12	10	13	Examples	Examples AA044081	486300 3'
		-	-	L				zk50g07.r1 Soares pregnant uterus NbHPU Homo sapiens cDNA clone
The state of the s								486300 5' similar to PIR. A40533 A40533 cAMP-dependent protein kinase
-							=	major membrane substrate
1 CATGAGGTCTTCAAT	H178129	4	2 0	09 (2	Examples X14787		Class A, Human mRNA for thrombospondin.
12 CATGAGGTGCGGGG	H178603	0	2 2		111	Examples R27738		yh64fi1.s1 Homo sapiens cDNA clone 134541 3'
		-	_					yj22f12.s1 Homo sapiens cDNA clone 149519 3' similar to SP ZK637.5
			-				H00276	CE00436 ARSA
			-					zm19d07.s1 Stratagene pancreas (#937208) Homo sapiens cDNA clone
13 CATGAGTATCTGGGA	H183787	6	<u>m</u>	15	73	Examples	Examples AA076235	526093 3'
		-	_				H13159	yj16c04.s1 Homo sapiens cDNA clone 148902 3'
		\vdash	-					2071c11.s1 Stratagene pancreas (#937208) Homo sapiens cDNA clone
							AA146632	592364 3*
3.1 CATGATACTTTAATT	H204740	-	0	3 18	6	Examples X80062	X80062	H.sapiens SA mRNA.
			-				169101	Human annexin V (ANX5) gene

-				-	-	H	_		X12454	Human mRNA for vascular anticoagulant
\dagger				\vdash	\vdash	-	_		M18366	Human placental anticoagulant protein (PAP) mRNA
+				\vdash	-	_	_		M21731	Human lipocortin-V mRNA, complete cds
+				-	_				J03745	Human endonexin II mRNA, complete cds
+			\vdash	-	_	_	_			GAMMA-INTERFERON-INDUCIBLE PROTEIN IP-30 PRECURSOR
3.5 C.	CATGATCAAGAATCC	H213518	ন	_	5 2	25	1 E	Examples 103909	103909	(HUMAN)
-					_	_				EST97384 Thymus II Homo sapiens cDNA 3' end similar to interferon,
									aa383911	gamma transducer 1
19 C	16 CATGATCAAGGGTGT	H213679	12	9	25 1	12 156		Examples U09953	U09953	Human ribosomal protein L9 mRNA
+			$ \cdot $	H	Н				U21138	Human ribosomal protein L9 mRNA, complete cds
									D14531	Human mRNA for human homologue of rat ribosomal protein
+			+	+	+	1	1			zm03a05.s1 Stratagene corneal stroma (#937222) Homo saniens cDNA
- <u>C</u>	CATGATCAAGTTCGA	H213751	0	7	- 00	3	10 E	xamples /	Examples AA063259	clone 513008 3'
9	400000000000000000000000000000000000000	U21612H	19		4	12	40—	Examples L42856	42856	RNA polymerase II transcription factor SIII p18 subunit mRNA
2 2	TO CATGATGAACTICG	H229502	-	0	1		4	Examples Z59242		H.sapiens CpG DNA, clone 13a10, reverse read cpg1
-			-							
1-2	US CAMBERCANA ABORD	H235531	2		12	2	22 B	Examples Z25820	225820	H.sapiens mRNA for mitochondrial dodecenoyl-CoA dehydrogenase
1-			-	\vdash	-		L		1.24774	Homo sapiens delta3, delta2-CoA-isomerase mRNA
12	CATGATGTCTTCGTT	H243676	0	0	-	0	14 E	Examples M84711	Vf84711	40S RIBOSOMAL PROTEIN S3A (HUMAN)
2 :	1) CATGATGTCTTTTCT	H243710	-	2	<u> </u>	14		Examples M62403	M62403	Human insulin-like growth factor binding protein 4
-				-						Human insulin-like growth factor binding protein 4 (IGFBP4) gene,
	KOO KAMO MOM KOM	UDAAART	+=	+	44		77	Examples 7,334.57		promoter and complete cus H semiens mts1 gene
7	CAIGAIGIGIAACGA	1	,	1				-		Human CAPL protein mRNA, complete cds
1 0	11 CATGCAACTTAAAGC	H270083	0	-	2 10		<u>—</u>	Examples N23207		yx70b09.s1 Homo sapiens cDNA clone 267065 3' similar to gb:L12350 THROMBOSPONDIN 2 PRECURSOR (HUMAN)
-				-						2125e11.s1 Soares ovary tumor NbHOT Homo sapiens cDNA clone 714188
5	15 CATGCACCTGTCCTT	H286424	0	4	2 10	-		xamples /	53	3' similar to gb:M33680 CD81 ANTIGEN (HUMAN)
-			+		1					CD81 antigen
C,	46 CATGCACTCAATAAA	H291889	0	6	7		19 E	Examples 1778203		Neurosin
-			\dashv	\dashv	_	_	4		U62801	protease M

							-		
17 CATGCAGCCTGGGGGC	H300971	. 0	- 0	- 0	0	01	Examples AA149942		2068d04.s1 Stratagene pancreas (#937208) Homo sapiens cDNA clone 592039 3' similar to TR:E218488 E218488 TRYPTASE
					 				zp66b09.11 Stratagene endothelial cell 937223 Homo sapiens cDNA clone 625145 5' similar to gb:M16937 HOMEOBOX PROTEIN HOX-B7
18 CATGCAGCGCGCCT	H301462	4	=	12	10	21 I	Examples AA187553		(HUMAN); contains element MER22 repetitive element
				-			-	M16937	Homeobox protein HOX-B7
19 CATGCAGGTTGTCCT	H307126	0	0	4	0	10	No Match		
SUICATGCAGTCTCTCAA	H309109	7	9	9		17 I	Examples U14972		Human ribosomal protein S10 mRNA
SI CATGCATCCCGTGAC	H316857	0	3	5	3	13 E	Examples U27293		Human leukotriene A4 hydrolase gene
			-	-	_		-		Human leukotriene A-4 hydrolase mRNA, complete cds
			+	-	_		-5	J02959	Human leukotriene A-4 hydrolase mRNA, complete cds
SYCATGCATTCCTCCTT	H325080	0	7	5	13	3	Examples X82434		H.sapjens mRNA for emerin
STOATGCCACCCCACC	H333138	3	1	17	18	2 F	Examples M88338		Human serum constituent protein (MSE55) mRNA
ST CATGCCAGTGGCCCG	H339606	23	=	37 2	22 5	S6 E	Examples U14971		Human ribosomal protein S9 mRNA
SSCATGCCATTTCTGG	H344031	0	7	9		10 E	Examples L01697		Homo sapiens alpha-1 type XV collagen mRNA
\$6 CATGCCCAAGCTAGC	H344691	61	∞	8	18 4	44 E	Examples X54079		Human mRNA for heat shock protein HSP27.
			-	-		_	2	ZZ3090	H.sapiens mRNA for 28 kDa heat shock protein
			-	_	_	_	15	X16477	Human mRNA fragment for estrogen-regulated 24k protein
			-	-	_		S	\$74571	estrogen receptor-related protein=27-kda heat shock protein
STUREGOODED	H347489	20	15	43	19 6	61 E	Examples X69392		H.sapiens mRNA for ribosomal protein L26.
			\vdash	-	_	_			Human ribosomal protein L26 (RPL26) gene
SS CATGCCCCTGCAGA	H350099	0	~	9	14 2	25 E	Examples U40434		Human mesothelin or CAK1 antigen precursor mRNA
			-	-	_				Human mRNA for pre-pro-megakaryocyte potentiating factor, complete
							니	D49441	cds.
SUCATGCCCGCATAGAT	H353481	0	9	0	8	11 E	Examples U12819		Human p16-INK4 (p16) gene
			\vdash	_	_		1	U38945	Human hypothetical 18.1 kDa protein (CDKN2A) mRNA
			-	_	_	_			MTS1=multiple tumor suppressor 1/cyclin-dependent kinase 4 inhibitor
							S	S69804	plé
			-	<u> -</u>		L	S	269822	CDK41=cyclin-dependent kinase 4 inhibitor
			-	-	_	_			tumor suppressor gene, P16/MITS1/CDKN2=cell cycle cycle negative
	`		\dashv	\dashv	_	1	S	S78535	regulator beta form
60]CATGCCTCCTGGG	H357867	∕ ∞	7	5 14		34 E	Examples 247319		H.sapiens mRNA for expressed sequence tag (clone 21fi7119)

							AA398406	zt60h12.s1 Soares testis NHT Homo sapiens cDNA clone 726791 3'
41 CATGCGGCCCTACC	H370034	4	4	4	2	Examples U21049	J21049	Human DD96 mRNA
AS CONTRACTOR AS	H387925	0	7	30	66	Examples X03212	₹03212	KERATIN, TYPE II CYTOSKELETAL 1
70					-		A A 187637	zp73f01.s1 Stratagene HeLa cell s3 937216 Homo sapiens cDNA clone 625849 31
		╀		+	+			zp35g11.s1 Stratagene muscle 937209 Homo sapiens cDNA clone 611492
63 CATECOTTTGAACAG	H392709	<u>~</u>	3	7	23	Examples AA176457	1A176457	3' similar to TR:G663269 G663269 BOLA
		-		-	-			zp35e11.s1 Stratagene muscle 937209 Homo sapiens cDNA clone 611468
						/	AA176541	3' similar to TR:G663269 G663269 BOLA.
64 CATGCGCCGACGATG	H415844	21 13	45	75	1	Examples X02492		Human interferon-inducible mRNA fragment
65 CATGCTCAACAGCAA	H475429	2	5 10	9	17	Examples T53402		ya88g05.s1 Homo sapiens cDNA clone 68792 31
								and a MMC analysis and HILLION HILLIAM SAN AND A SAN FELL
							W69493	204/goo.st Somes tetal meat Pormats w mound saptems Court Cooles 343838 3' similar to PIR:S24168 S24168 hypothetical protein - human
	1475478	\rac{1}{-}	2	23	┾	Examples X13916	T	Human mRNA for LDL-receptor related protein
OD CATECOCOCOC	H493576	- -		000	180	Examples X80335	X80335	H.sapiens (24) Ferritin H pseudogene.
COLUMN CONTRACTOR CONT	H494454	1	4	21	13	Examples X04828	(04828	Human mRNA for G(f) protein alpha-subunit
ADOTATE CONTROLL CANADA CONTROLL CANADA CONTROLL CONTROL	H49887	16 30	28	30	4	Examples U14966	714966	Human ribosomal protein L5 mRNA
TO CAMECTECT GRANGER	H499247	-	3	13	13	Examples T90665	190665	yd41g08.s1 Homo sapiens cDNA clone 110846 3'
100000000000000000000000000000000000000		-		-	-			EST43791 Fetal brain I Homo sapiens cDNA 3' end similar to steroid
							AA338799	hormone receptor NERR1
		-		-			H97236	yv98b06.s1 Homo sapiens cDNA clone 250739 3'
71 CATGCTGGCGCGAT	H501337	0	4	0	10	Examples C14084	314084	Human fetal brain cDNA 3'-end GEN-018D10
7) CATGCTTCCAGCTAA	H513181	64 23	36	53	104	Examples D00017	200017	Human lipocortin II mRNA
73 CATGOTTGCCT	H514022	0	4	68	7	Examples Z19574	219574	H.sapiens gene for cytokeratin 17.
		_		-	-		X62571	H.sapiens mRNA for keratin-related protein
		_		-	-	^	X05803	Human radiated keratinocyte mRNA 266
711CATGCTTTCCTT	H522198	0	7	16	4	Examples X79067	£79067	H.sapiens ERF-1 mRNA 3' end.
75 CATGGAAAAAAAA	H524289	7 14	21	76	37	Examples X51779	451779	Human mRNA containing an Alu repeat
		_		-	-	<u> </u>	X82240	H.sapiens mRNA for Teell leukemia/lymphoma 1
76 CATGGAAACAAGATG	H525348	4	7 14	8	22	Examples	V00572	Human mRNA encoding phosphoglycerate kinase.
		_				1	D29018	Human keratinocyte cDNA, clone 001
						1		Human phosphoglycerate kinase (pgk) mRNA
77 CATGGAAATACAGTT	H527436	49 3	35 10 100	100	36	Examples X05344		Human mRNA for cathepsin D
יייייייייייייייייייייייייייייייייייייי				-				

		-	-			N N	M11233	Human cathepsin D mRNA, complete cds
		+	+	1	+			vol42/ft/3 s1 Home samens cDNA clone 110909 3' similar to SP:R151.9
. CATGGAAATGATGAG	H527929	4	7 5	14	26	Examples T90296	90296	CE00827
						¥	AA320942	EST23523 Adipose tissue, brown Homo sapiens cDNA 3' end
		+	-					zp64f07.s1 Stratagene endothelial cell 937223 Homo sapiens cDNA clone
	9£7££5H	٠,	7 16	9	78	Examples AA 181811	٠,	624997 3'
CAlecandation		+	-	1		-		z106c06.s1 Soares pregnant uterus NoHPU Homo sapiens cDNA clone
						_	AA148508	491530 3' similar to WP:ZK652.2 CE00448
CATTTTT CANDED	H540621	9	3 10	6	28	Examples L21950	21950	Human peripheral benzodiazepine receptor related mRNA
NO CALGORATITIONS		+	-		_	2	M36035	Human peripheral benzodiazepine receptor (hpbs) mRNA
- I Charles and Adda Adda Adda Adda Adda Adda Adda	H540673	-	2 10	3	17	No Match		
CATCONCONTRACTOR	H545152	0	0	Ξ	7	Examples U19718		Human microfibril-associated glycoprotein (MFAP2).
TO TO A CONTRACTOR OF THE PARTY	H545430	0	3	22	18	Examples M75165		H.sapiens epithelial tropomyosin (TM1) mRNA
ייייין פפטרביטיפיייי		┝	-		<u> </u>	W	M12125	Human fibroblast muscle-type tropomyosin mRNA
		+	\vdash			Σ	M74817	Human tropomyosin-1 (TM-beta) mRNA, complete cds
Jestanonanoma	H546059	2	5	19	2	Examples M74092	174092	Human cyclin mRNA
CAI GENCECCONTROCT	H546710	31	36 20	12	3	Examples L37033	37033	Homo sapiens FK-506 binding protein homologue
יאן פפארברבן פרבבן		1	-					2b37g02.s1 Soares parathyroid tumor NbHPA Homo sapiens cDNA clone
	H548062	-6	=	Ξ		Examples N90046	90046	305810 3'
A COLORANGE DE CECE		\vdash	-					zl06a10.s1 Soares pregnant uterus NbHPU Homo sapiens cDNA clone
-						Y	AA115048	491514 3'
004000000000000000000000000000000000000	H551315	<u>_</u>	4 5	32	3	Examples M63193	163193	Human platelet-derived endothelial cell growth factor
CALCACACACACACACACACACACACACACACACACACA	H554876	-	4	0	4	Examples M61764	161764	Human gamma-tubulin mRNA,
AN CATGOAGACTTTGC	H559615	0	0	7	2	Examples D17793	17793	Human mRNA (HA1753) for ORF
SECTION CONTROL OF COLOR	HS60056	0	5 8	32	Ξ	Examples S68252	68252	TIMP-1≈metalloproteinase inhibitor
200000000000000000000000000000000000000		\vdash	-			X	X02598	EPA glycoprotein (erythroid-potentiating activity)
		-	\vdash			×	X03124	tissue inhibitor of metalloproteinase 2
"I CATGGAGCAGGATGA	H561807	0	0		12	No Match		
ひしかもじ なししし なじし せんじ でこ	H567486	-=		4	13	Examples AA214523	A214523	2189c01.s1 Soares NbHTGBC Homo sapiens cDNA clone 682848 3'
V. LAI GGAGGGAGTTCC		+	-		 -	Z	N30324	yw75d01.s1 Homo sapiens cDNA clone 258049 3'
us carecaserceses	H570787	0	0 2	_	01	Examples X70070		H.sapiens mRNA for neurotensin receptor.
CHICAMENT TO CO.	H577656	-	3	0	2	Examples H57673		yr27a10.s1 Homo sapiens cDNA clone 206490 3'
94 CATGGAGTIALGILG	1	1						

		-	_	_		•		
		+	1	+	\dagger		M73240	Human (clone SF2) heratacyte growth factor (HGF)
TOUTON A A ACTUAL	H655547	18 13	3	10/2	+	Examples X02920	X02920	Human mRNA for alpha 1-antitrypsin carboxyterminal, 0
100100000000000000000000000000000000000		-		\vdash	-		X01683	Human mRNA for alpha 1-antitrypsin
		\vdash		-	-		V00496	Human messenger RNA for alpha-1-antitrypsin
		-		\vdash			100067	Human alpha-1 antitrypsin gene, 3' end
		-		\vdash	-			zi22b01.s1 Soares pregnant uterus NbHPU Homo sapiens cDNA clone
	H658059	0	4	9	16	Examples	Examples AA127040	502633 3'
200000000000000000000000000000000000000		-			-			zd86f06.s1 Soares fetal heart NbHH19W Homo sapiens cDNA clone
							W81387	347555 3'
		-		-	-		H45477	yo72h08.s1 Homo sapiens cDNA clone 183519 3'
	H666943	9	9	2	32	Examples D26598	D26598	Human mRNA for proteasome subunit HsC10-II. , 0
CAT GGGAGT COTTON	H667367		0	-	2	Examples N74310	N74310	za78c01.s1 Homo sapiens cDNA clone 298656 3'
157616161616171				-	\vdash		H92750	yt92e01.s1 Homo sapiens cDNA clone 231768 3'
				-			174084	sen 2272 Homo saniens cDNA clope ssb4HB3MA(extended-ft-6) 3'
	2571730	` -	15	~	=	Examples X17567	X17567	H. sapiens RNA for snRNP protein B
CATGGGATTGTCTGG	CCTIVAL	,		+	+		M34081	Human small nuclear ribonucleoprotein particle SmB
	H677330	0	0	6	22	Examples M69054	M69054	Human insulin-like growth factor binding protein 6, 0
200000000000000000000000000000000000000		-		-	-		M62402	Human insulin-like growth factor binding protein 6
540000000000000000000000000000000000000	H677753	0	4	1	14	Examples N74323	N74323	za78d08.s1 Homo sapiens cDNA clone 298671 3'
יאו פפררכיז כז פאס		+	L	-	\vdash		H46766	yo18f08.s1 Homo sapiens cDNA clone 178311 3'
				-	-		H41102	yn88a08.s1 Homo sapiens cDNA clone 175478 3'
		-		-	 -			zm84b09.s1 Stratagene ovarian cancer (#937219) Homo sapiens cDNA
554345543555457	H686815	0	3	13	77	Examples	Examples AA074777	clone 544601 3'
		\vdash		-	-			zm04a04.51 Stratagene comeal stroma (#937222) Homo sapiens cDNA
							AA062735	clone 513102 3'
		-		\vdash	-			zm63f12.s1 Stratagene fibroblast (#937212) Homo sapiens cDNA clone
							AA112905	530351 3'
117 CATGGGGAAGCAGAT	H688713	25	7 9	0	72	No Match		
18 CATCCCCAAGGGGTGG	H690863	7	3 1	16	7	No Match		
19 COTTOGGGAGGTAGCA	H690890	-	-	14	-	No Match		
130 CATGGGGGGATCTCTT	H693112	-	3	39	77	Examples V00523	V00523	Human mRNA for histocompatibility antigen HLA-DR
		-		-	-		X00274	Human gene for HLA-DR alpha heavy chain a class II
			1	-				III III A DD Alaka abain ambhild

		r	-	-	L	L	<u>S</u>	100202	human hla-dr heavy chain gene; 3' flank
VI CATCCCTCCCCAGAT	H715401	-	4	01	2	14 E	Examples U18009		Human chromosome 17q21 mRNA clone LF113.
100000000000000000000000000000000000000		T	-	\vdash			13	T33413	EST57778 Homo sapiens cDNA 3' end similar to None
		<u> </u>	-	_	_	L	T3.	T33339	EST57474 Homo sapiens cDNA 3' end similar to None
CATGGTACTGTAGCA	H728778	6	3	-	16	30 E	Examples M59911		Human integrin alpha-3 chain mRNA
171 CATGGTACTGTGGCT	H728810	23	101	16	'	50 E	Examples X87689		H.sapiens mRNA for putative p64 CLCP protein
CATEGACABATTTC	H737344	0	0	0	101	1 E	Examples L12350		Human thrombospondin 2 (THBS2) mRNA
174 CATGGTCTGGGGCTT	H752296	25	35 4	45 76		29 E	Examples D21261		Human mRNA (HA1756) for ORF
171		\vdash	-	_		_	DZ		Human keratinocyte cDNA, clone 686
SAS CATCHERON OF SCI	H752521	0	2	7 12	2	2 E	Examples H51290		yp07a05.s1 Homo sapiens cDNA clone 186704 3'
			-	_	_	L	NZC		yx44g12.s1 Homo sapiens cDNA clone 264646 3'
		╁	-	_	_	_			2076e09.s1 Stratagene pancreas (#937208) Homo sapiens cDNA clone
							AA	AA158271	592840 3'
131 CATGETCTGTGCAGG	H752531	0	0	0		13 K	No Match		
178 CATGGTCTTGAAGCC	H753162	0	-	7		10 N	No Match		
130 CATGGTGAAGGCAGT	H754323	25	14	42 15	_	89 E	camples X8		Class C, H.sapiens RPS3a gene
130 CATGGTGAATGACGG	H754567	0	2	00		10 E	Examples X08058		GLUTATHIONE S-TRANSFERASE P (HUMAN)
131 CATGGGGGGGGGG	H760361	0	<u>~</u>	2 11		25 E	Examples X51439		Human mRNA for serum amyloid A (SAA) protein
133 Characterage BAA	H761481	7	5	13	L		Examples U15008		Human SnRNP core protein Sm D2 mRNA
133 CATEGING AGGGCAC	H762533	-	-	8	3	34 E	Examples U62800		Cystatin M (CST6)
13 1 CATCOTOCTACAGGA	H765003	7	171	15 39			Examples H46430		yo12h12.s1 Homo sapiens cDNA clone 177767 3'
100000000000000000000000000000000000000		1	-	-	L	_			zf13a06.s1 Soares fetal heart NbHH19W Homo sapiens cDNA clone
							¥¥	AA047563	376786 3'
			_	-	L	_			zo13f02.s1 Stratagene colon (#937204) Homo sapiens cDNA clone 586779
							AA	10	31
135 CATGGTTCACTGCAG	H774629	0	7	1 13		3	Examples X59288		H.sapiens gene for intercellular adhesion molecule
				_		_	M2		Human major group rhinovirus receptor (HRV) mRNA
		-	-	_	_		103	103132	Human intercellular adhesion molecule-1 (ICAM-1)
			-	_	_	_	MS		Human cell surface glycoprotein P3.58 mRNA
1 36 CATGGTTGTCTTTGG	H781823	F	-	6 30		24 E	Examples K02765		Human complement component C3 mRNA, alpha and beta
117 CATGGTTGTGTTAA	H782013 178		11011	14 340	0 139		Examples MI7987		Human beta-2-microglobulin gene
138 CATGGTTTAAATCGA	H782391	=	9	12	4	14 E	Examples D00760		Human mRNA for proteasome subunit HC3
CAREELUCARELES	H797169	-	0	9		12 臣	Examples X57025		INSULIN-LIKE GROWTH FACTOR IA PRECURSOR (HUMAN)
CALCIA CALGIA MATATAGEN	H802793	0	77	_	7		No Match		
170 CW101W111100		1	-	j					

CATGTAATTTTGGAT	H802793	-	-			No Match		
11 CATCTACTACTORT	H806901	-	4	m	7	Examples X85373		H.sapiens mRNA for Sm protein G
TO THE STREET OF THE STREET	H808370	0	1 4	0	2	No Match		
11 CATGTACCCTTCTAT	H808925	0	0	=	7	No Match		
11 CATCTACGAAAGTAA	H827437	-	0	S.	77	Examples 102931	102931	Human placental tissue factor (two forms) mRNA
100000000000000000000000000000000000000		\vdash	_		Γ		M16553	Human tissue factor mRNA, complete cds
		+	L				M27436	Human tissue factor gene, complete cds
18 CarcabecerrentA	H831416	49	61 61	8	130	Examples X64899	X64899	H.sapiens mRNA homologous to mouse P21 mRNA.
100000000000000000000000000000000000000		-	_				X16064	Human mRNA for translationally controlled tumor protein
		-					L13806	Homo sapiens (clone 04) translationally controlled tumor protein
CATGTATATTTTCTC	H839672	-	0 3	∞	19	Examples M98479		Human transglutaminase mRNA
1. CATGRATTTCTGCC	H851834	0	1 2	16	٣	Examples D12149		Human HepG2 3'-directed Mbol cDNA, clone s247
18 CATCTCACAAGCAAA	H856209	10 2	28 27	24	48	Examples X80909		H.sapiens alpha NAC mRNA
LOCATGLCCAAATCGAT	H868569	0	1 0	43	17	Examples X56134	X56134	Human mRNA for vimentin.
		_					Z19554	H.sapiens vimentin gene
		-	<u> </u>				M14144	Human vimentin gene, complete cds
		-	_			-	M25246	Human vimentin (HuVim3) mRNA, 3' end
SHETCCACTGGCCT	H870310	0	1	12	2	Examples N92906	N92906	zb57a08.s1 Homo sapiens cDNA clone 307670 3'
						<u> </u>	T17488	NIB978 Normalized infant brain, Bento Soares Homo sapiens cDNA 3'end
		+					AA349906	EST56900 Infant brain Homo sapiens cDNA 3' end
STOP TO TOTAL CATE OF THE	H871920	9	6 10	25	150	Examples X67016	X67016	H.sapiens mRNA for amphiglycan
		-	_				D13292	Human mRNA for ryudocan core protein
SOCATGTCGTCTTTATC	090668H	7	5 15	E	69	Examples M77233	M77233	Human ribosomal protein S7 mRNA
SATGTCTCTGATGCT	H908858	-	5 2	46	19	Examples S48568	\$48568	tissue inhibitor of metalloproteinase 2 (3'-end region)
		-	1					
1511 CATGTCTTGTBACTG	H916232	0	4 3	-	E	Examples N71680		yz93b03.s1 Homo sapiens cDNA clone 290573 3'
155 CATGEOTTGEGGATA	H916372	14 2	22 15	8	45	Examples X03083		Human lactate dehydrogenase-A gene
		-	_					Human mRNA for lactate dehydrogenase-A
	,	-	_				X02153	Human pseudogene for lactate dehydrogenase-A
SOCATGTGAAGTCACTG	H920392		1 6	0	16	No Match		
	H920525	0	1 3	· v	==	Examples X07979		CTGTGG, Class A, Human mRNA for fibronectin receptor beta subunit.

				-	-				zk05h07,s1 Soares pregnant uterus NbHPU Homo sapiens cDNA clone
2	TESTITEM ASTORACIONS	H932731	٥	∞	3	11 1	12 Examp	Examples AA027860	469693 3'
3	SO CATOTOCOLORIO	H938876	-	<u></u>	7	3	16 Exam	Examples M25753	G2/MITOTIC-SPECIFIC CYCLIN B1 (HUMAN)
:				-				T60151	yc22c04.s1 Homo sapiens cDNA clone 81414 3'
			ŀ	H	-			R67969	yi29g08.s1 Homo sapiens cDNA clone 140702 3'
						······			2091f03.s1 Stratagene ovarian cancer (#937219) Homo sapiens cDNA
100	ななないとしていりませんという	H939841	=======================================	13	3 13		43 Exam	Examples AA169614	GIONE 594269 5 SIMILIA TO SWINDAL, HUMAN PROTOS INCUFRICAL GELATINASE-ASSOCIATED LIPOCALIN PRECURSOR
8	33.000000000000000000000000000000000000	1020840	"	-	=		19 Exame	Examples N79823	zb15d08.s1 Homo sapiens cDNA clone 302127 3' similar to SW:NGAL_HUMAN P80188 NEUTROPHIL GELATINASE- ASSOCIATED LIPOCALIN PRECURSOR
3	161 CATGTGCCCTCAGAA	11222047	1	+					
									zm90h04.s1 Stratagene ovarian cancer (#937219) Homo sapiens cDNA clone 545239 3' similar to SW:NGAL_HUMAN P80188 NEUTROPHIL
162	162 CATGTGCCCTCAGGA	H939851	17	3	10 25	83		Examples AA075896	GELATINASE-ASSOCIATED LIPOCALIN PRECURSOR
162	162 CATGTGCCCTCAGGC	H920392	1	+	4	1	No Match	Itch	-181-07 cl Stratement Colon (#037704) Home saniens CDNA clone 511044
		22011011	c	r			12 Examt	Examples AA 100279	216150 /.S1 Suatagene contr. (#25/204) Monto September 1.1.1. 1.1.1.1.1.1.1.1.1.1.1.1.1.1.1.1
9	163 CATGI GCCI TACLI I	H944038	1	1	2 17	L		tch	
Į.	164 CATGTGCGCT GGCCC	2001	1	+	+	_			2k10a01.s1 Soares pregnant uterus NbHPU Homo sapiens cDNA clone
		H949560	7	9	9	1	16 Examp	Examples AA029262	470088 31
	22120112010			+	-	_			yv66e10.s1 Soares fetal liver spleen INFLS Homo sapiens cDNA clone
								N54281	247722 3'
				}				A A 1 14075	zn76c02.s1 Stratagene NT2 neuronal precursor 937230 Homo saptens cDNA clone 564098 3'
		H953251	8	15	7 22	1	48 Exam	Examples L76200	Homo sapiens guanylate kinase (GUK1) mRNA
9 5	CATGLGGAGIGGAGG	H955723	0	m	3 37		_	Examples X00570	Human mRNA for precursor of apolipoprotein Cl
3	CAT GOOD ON THE COLUMN	H962086	13	15	13 76		27 Exam	Examples L16510	Homo sapiens cathepsin B mRNA
				-	-	_		M14221	Human cathepsin B proteinase mRNA, complete cds
1031	TOOODS ASTROPHENT OF	H975446	6	m	3 22		8 Exam	Examples L35240	Human enigma gene
	TO CATGLETIC TAPATE	H976644	∞	21	26 18		50 Exam	Examples L38941	Homo sapiens ribosomal protein L34 (RPL34) mRNA
	TICATGTGTGTGTTTGT	H978687	9	7	16 25		15 Exam	Examples X03473	(Human gene for histone H1(0).
		400000	-	-	1 21		1 Exam	Examples AA034505	zk23g08.si Soares pregnant uterus NbHPU Homo sapiens cDNA clone 471422 3'
	1-2)CATGTTATGGATCIC	117777		+	-1				

						AA235464	31
						AA037024	zk30c10.s1 Soares pregnant uterus NbHPU Homo sapiens cDNA clone 472050 31
- STOTTCATTGTAGA	H1003443 0	1	01	3	Examples H53629	H53629	yu38d04.s1 Homo sapiens cDNA clone 236071 3'
						T06706	EST04595 Homo sapiens cDNA clone HFBDX32
						T16635	NIB1599 Normalized infant brain, Bento Soares Homo sapiens cDNA 3'end similar to EST04595 H. sapiens cDNA clone HFBDX32
CATGTTCTGTGAATC	H1014660 3	4	3 24			Examples AA026678	2e97h02.s1 Soares fetal heart NbHH19W Homo sapiens cDNA clone 366963 3'
						AA280283	205a03.s1 Soares NbHTGBC Homo sapiens cDNA clone 712204 3'
			-			H10141	ym05a09.s1 Homo sapiens cDNA clone 46675 3'
379775775775	H1021276 0	0	0	8 17		X66029	H.sapiens mRNA for tyrosine kinase receptor.
- aretrecteactr	H1023520 1	N.	1 33		Examples X15880	X15880	Human mRNA for collagen VI alpha-1
			-	_		X72414	H.sapiens gene for glutaminyl-tRNA synthetase
			_				zk73h10.s1 Soares pregnant uterus NbHPU Homo sapiens cDNA clone
CATGGAGATCTC	H1024568 4	Ξ	16 10	24		Examples AA044568	488515 3'
						N71899	yz36b07.s1 Homo sapiens cDNA clone 285109 3'
						AA400793	2171g03.s1 Soares testis NHT Homo sapiens cDNA clone 727828 3'
TA PATETTEGGGTTTCC	H1026814 202	75	84 235	369	Examples	X80336	H.sapiens (5) Ferritin H pseudogene.
			_			X00318	Human mRNA for apofertitin H chain type
			-	_		X03488	Human apoferritin H gene exons 2-4
			_			M97164	Human ferritin heavy chain mRNA, complete cds
			_			L20941	Human ferritin heavy chain mRNA, complete cds
TOCATGTTGGTGAAGGA	H1027595 98	98 106	17 183	3 107	Examples X02493	X02493	Human interferon-inducible mRNA (cDNA 6-26).
			-			M11948	Human promyelocytic leukemia cell mRNA
			_			M17733	Human thymosin beta-4 mRNA, complete cds
AN CAMETTIC COTCADA	H1037777 0	-	0 13	3	Examples N78832	N78832	2b17a08.s1 Homo sapiens cDNA clone 302294 3'
			-	_			2t33d02.s1 Soares ovary tumor NbHOT Homo sapiens cDNA clone 724131
	,					AA411095	31
			_				zd84g11.s1 Soares fetal heart NbHH19W Homo sapiens cDNA clone
				_		11101100	347306 21

	1200000011	10	1	F	12	110000001 of 21 21 71 Example NO. 0171	Human brain tune clathrin light chain a mRNA
SILCATGITICCTICCIT	Q67850IH	5	7	_	-	Examples intend	minian oranic gpc comming in Sur Caram a mice com
			\vdash	-	-	M20472	Human lymphocyte clathrin light-chain A mRNA
INTERTEGRACETT	H1041504 2 0 0 16	2 0	0	91	-	Examples X78947	H.sapiens mRNA for connective tissue growth factor
			-	-	\vdash	U14750	Human connective tissue growth factor mRNA
IST CATGTTTGTTAAAA	H1044225		-	-	-	H06492	y178c08.s1 Homo sapiens cDNA clone 44273 3'
			-	-	\vdash	T35952	EST94173 Homo sapiens cDNA 3' end similar to None
			-	-	-	AA253218	AA253218 zr53g10.s1 Soares NhHMPu S1 Homo sapiens cDNA clone 667170 3'

Table 5 - Transcripts increased in pancreas and colorectal cancer

SAGE tag that were elevated in both in coloreactal and pancreatic tumor, and are likely to be specific for tumor in general.

Tag_Sequence Tag_Number Accession Description De					
C -950498 M10629 G -294155 U42376 U56145 T (A) -243747 J03040 M25746 C -610466 X53416 T -229106 X02761 T -229106 X02761 C -760291 X58536 G -760291 X58536 C -760291 X58536 C -760291 X58536 C -76920 M77349 C -76920 M77349 C -589267 X585379 T -8589267 X55928 T -8589267 X55989 C -884181 X15804 C -884181 X15804 C -884181 X15804 C -884181 X15804 C -673954 X17620 M19045 T -53129 U62962 A -53129 U62962 A -1048113 D16891	Tag	Segmence		Tag Number Accessic	
G294155 U42376 T (A) -243747 J03040 C -610466 X53416 T229106 X02761 T229106 X02761 C -760291 X58536 G -76231 M95787 C -76231 M95787 C -76920 M77349 C -589267 X53219 C T -8582 X57351 T -8582 X57351 T -8582 X57351 C T -615821 D80012 T -241665 M74090 M19045 C -673954 X17620 A -1048113 D16891 T -53129 U62962 A -1048113 D16891	1 CATE	TEGABATGAC	U	-950498 M10629	Human alpha-1 collagen gene,
T(A) -243747 J03040 T -229106 X53416 T -229106 X02761 T -229106 X02761 K00799 C -760291 X58536 A -760291 X58536 A -76920 M7349 C -589267 X53279 C -8892 X53279 C C -8892 X53219 T -8582 X57351 T -8582 X57351 T -615821 D80012 T -241665 M74090 C - 894113 D16891 M9945 A -53129 U62962 A -1048113 D16891	2 CATC	CACTTCAAGG	U	-294155 042376	Θ
T(A) -243747 J03040 C -610466 X53416 T -229106 X02761 T -229106 X02761 K00799 C -760291 X58536 G -76231 M95787 C -769020 M77349 C -589267 X53279 T -85822 X57321 T -85822 X57351 T -85821 X57359 C T -884181 X15804 C T -241665 M74090 C T -241665 M74090 C T -241665 M74090 A -1048113 D16891 T -241665 M78093 A -1048113 D16891 T -302741 X53743	2			056145	Human thymic shared antigen-1/stem cell antigen-2
C -610466 X53416 T -229106 X02761 C -610466 X53416 C -760291 X58536 G -76231 M95787 C -76231 M95787 C -76231 M95787 T -889267 X53279 C -884181 X15804 C -884181 X15804 C -884181 X15804 C -673954 X17620 A -53129 U62962 A -1048113 D16891 T -51329 U62962			T(A)	-243747 J03040	Human SPARC/osteonectin mRNA, complete cds.
C -610466 X53416 T -229106 X02761 C -760291 X58536 G -76231 M95787 G -76231 M95787 C -769020 M77349 C -589267 X53279 C -884181 X15804 C -884181 X15804 C T -615821 D80012 T -241665 M74090 C -884181 X15804 C -884181 X15804 C -873954 X17620 M19045 C -673954 X17620 A -1048113 D16891 T -302741 X53743				M25746	- 1
T -229106 X02761 C -760291 X58536 G -76231 M95787 G -76231 M95787 C -769020 M77349 C -589267 X53279 C -589267 X53279 C -884181 X15804 C -884181 X15804 C -884181 X15804 C - 873954 X17620 M -53129 U62962 A -53129 U62962	4 CATG	GCCCAAGGAC	O	-610466 X53416	amin)
C -760291 X58536 G -76221 M26432 G -76231 M95787 C -589267 X53279 C -589267 X53279 C -589267 X53279 C -884181 X15804 C -884181 X15804 C -884181 X15804 C - 673954 X17620 M 19045 C - 673954 X17620 A -53129 U62962 A -53129 U62963 A - 1048113 D168963	SCATG	ATCTTGTTAC	F	-229106 X02761	Human mRNA for fibronectin (FN precursor).
C				K00799	human fibronectin (fn) 3' coding region and flank,
G -76231 M95787 G -76231 M95787 C -589267 X53279 C -589267 X53279 T -8582 X57351 T -8582 X57351 C -884181 X15804 C,T -515821 D80012 T -241665 M74090 C -673954 X17620 A -53129 U62962 A -53129 U62962 A -1048113 D16891 T -302741 X53743	6 CATG	GTGCGCTGAG	U	-760291 X58536	Human mRMA for HLA class I locus C heavy chain.
G -76231 M95787 A -769020 M77349 C -589267 X53279 X55958 J04948 T -85822 X57351 C -884181 X15804 C,T -515821 D80012 T -241665 M74090 C -673954 X17620 A -1048113 D16891 A -1048113 D16891 T -53129 U62962	2			M26432	Human MHC class I HLA-C.1 gene, complete cds.
A -769020 M77349 C -589267 X53279 X55958 J04948 T -85822 X57351 C -884181 X15804 C,T -515821 D80012 T -241665 M74090 C -673954 X17620 A -1048113 D16891 T -53129 U62962 A -1048113 D16891 T -302741 X53743			9	-76231 M95787	- [
A -769020 M77349 C -589267 X53279 X55958 J04948 T -85882 X57351 C -884181 X15804 C,T -515821 D80012 T -241665 M74090 C -673954 X17620 A -1048113 D16891 T -302741 X53743				M83106	Human SM22 mRNA, 5' end.
C -589267 X53279 X55958 J04948 T -85882 X57351 C -884181 X15804 C,T -515821 D80012 T -241665 M74090 C -673954 X17620 A -1048113 D16891 T -302741 X53743	8 CATG	GTGTGTTTGT	A	-769020 M77349	Human transforming growth factor-beta induced gene
T -85882 X57351 C -884181 X15804 C,T -515821 D80012 T -241665 M74090 C -673954 X17620 C -673954 X17620 A -1048113 D16891 T -302741 X53743	STAT 6	GATTTCTCAG	U	-589267 X53279	Human mRNA for placental-like alkaline phosphatase
T -85882 X57351 C -884181 X15804 C,T -515821 D80012 T -241665 M74090 C -673954 X17620 C -673954 X17620 A -1048113 D16891 T -302741 X53743				X55958	H. sapiens mRNA for alkaline phosphatase.
T -85882 X57351 X02490 C,T -515821 D80012 T -241665 M74090 C -673954 X17620 C -673954 X17588 A -1048113 D16891 T -302741 X53743				304948	Human alkaline phosphatase (ALP-1) mRNA, complete
C -884181 X15804 C, T -515821 D80012 T -241665 M74090 C -673954 X17620 C -673954 X17620 A -53129 U62962 A -1048113 D16891 T -302741 X53743	101CATG	ACCATTCTGC	E	-85882 X57351	Human 1-8D gene from interferon-inducible gene fam
C, T -515821 D80012 T -515821 D80012 T -241665 M74090 C -673954 X17620 C -673954 X17620 A -53129 U62962 A -1048113 D16891 T -302741 X53743	2112			X02490	Human interferon-inducible mRNA (cDNA 1-8).
C,T -515821 D80012 T -241665 M74090 C -673954 X17620 C -673954 X17620 A -53129 U62962 A -1048113 D16891 T -302741 X53743	11 CATG	TCCTTCTCCA	U	-884181 X15804	Human mRNA for alpha-actinin.
T -241665 M74090 D03801 C -673954 X17620 X75598 A -53129 U62962 A -1048113 D16891 T -302741 X53743	12 CATG	CTTCTGTGTA	C, T	-515821 080012	Human mRNA for KIAA0190 protein.
C -673954 X17620 X75598 A -53129 U62962 A -1048113 D16891 T -302741 X53743	13 CATG	ATGTAAAAA	T	-241665M74090	Human TB2 gene mRNA, 3' end.
C -673954 X17620 X75598 A -53129 U62962 A -1048113 D16891 T -302741 X53743				103801	Human lysozyme mRNA, complete cds with an Alu repe
C -673954 X17620 X75598 A -53129 U62962 A -1048113 D16891 T -302741 X53743				M19045	Human lysozyme mRNA, complete cds.
A -53129 U62962 Human Int-6 mRNA, complete A -1048113 D16891 Human HepG2 3' region cDNA, T -302741 X53743 H.sapiens mRNA for fibulin-	14 CATG	GGCAGAGGAC	O	-673954 X17620	Human mRNA for Nm23 protein, involved in developme
A -53129 U62962 Human Int-6 mRNA, complete A -1048113 D16891 Human HepG2 3' region cDNA, T -302741 X53743 H.sapiens mRNA for fibulin-				X75598	H.sapiens nm23Hl gene.
A -1048113 D16891 Human HepG2 3' region cDNA, T -302741 X53743 H.sapiens mRNA for fibulin-	15 CATG	AATATTGAGA	A	-53129 062962	Human Int-6 mRNA, complete cds.
T -302741 X53743	16 CATG	TTTTGATAA	A	-1048113D16891	Human RepG2 3' region cDNA, clone hmd2c11.
	17 CATG	CAGCTGGCCA	Į.	-302741 X53743	H.sapiens mRNA for fibulin-1 C.

-774461 X00497 Human mRNA for HLA-DR antigens associated invarian	M13560 Human Ia-associated invariant gamma-chain gene, ex	-2056 Y00345 Human mRNA for polyA binding protein.	58533 M61831 Human S-adenosylhomocysteine hydrolase (AHCY) mRNA	M61832 Human S-adenosylhomocysteine hydrolase (AHCY) mRNA	918273X16934 Human hB23 gene for B23 nucleophosmin.	M28699 Homo sapiens nucleolar phosphoprotein B23 (NPM1) m	M23613 Human nucleophosmin mRNA, complete cds.	M26697 Human nucleolar protein (B23) mRNA, complete cds.	-998030 M24194 Human MHC protein homologous to chicken B complex	-274492 D23661 Human mRNA for ribosomal protein L37, complete cds	L11567 Homo sapiens ribosomal protein L37 mRNA, complete	-155632 D83174 Human mRNA for collagen binding protein 2.	terferon-inducible	hoprotein	J05068 human transcobalamin I mRNA, complete cds.	-398663 M12529 Human apolipoprotein E mRNA, complete cds.	K00396 Human apolipoprotein E (epsilon 2 and 3 alleles) m	-298495 X56998 Human UDA52 adrenal mRNA for ubiquitin-52 amino ac	X56999 Human UbA52 placental mRNA for ubiquitin-52 amino	501287 X07491 Human DNA inserts showing sperm-specific hypomethy	M91670 Human ubiquitin carrier protein (E2-EPF) mRNA, com	-256497 L14272 Human prohibitin (PHB) gene, exons 1-7.	S85655 prohibitin [human, mRNA, 1043 nt].	-765573 U62435 Human nicotinic acetylcholine receptor alpha6 subu	U68041 Human breast and ovarian cancer susceptibility pro	-883029M24398 Human parathymosin mRNA, complete cds.	-125661 X58965 H.sapiens RNA for nm23-H2 gene.	M36981 Human putative NDP kinase (nm23-H2S) mRNA, complet	L16785 Homo sapiens c-myc transcription factor (puf) mRNA	-33331 U02032 Human ribosomal protein L23a mRNA, partial cds.	037230 Human ribosomal protein L23a mRNA, complete cds.	
G -774461		T -2056	G -58533		C -918273				T -998030	T -274492		6 -155632	C97078	A -1000193		C -398663		T -298495		C -501287		A -256497		C -765573		T -883029	T -125661			A -33331		
I BICATG GTTCACATTA		19 CATG AAAAGAAACT	20 CATG AATGCAGGCA		21 CATG TGAAATAAAA				22 CATG TTATGGGATC	CATG CAAT		24 CATG AGCCTITGIT		26 CATG TTCAATAAAA		27 CATG CGACCCCACG		28 CATG CAGATCITIG		29 CATG CTGGCGAGCG		30 CATG ATTGGCTTAA		31 CATG GTGGTGGACA		32 CATG TCCTGCCCCA	CATG			34 CATG AAGAAGATAG		

	L13799	Homo sapiens (clone 01) liver expressed protein mR
25 CATC ACATCATCA T	-79065 L06505	Human ribosomal protein L12 mRNA, complete cds.
CATG CTGTTGGTGA	-507577 D14530	Human homolog of yeast ribosomal protein S28, comp
ATTA	-249854 X57959	H.sapiens mRNA for ribosomal protein L7.
	X57958	H.sapiens mRNA for ribosomal protein L7.
	X52967	Human mRNA for ribosomal protein L7.
	116558	Human ribosomal protein L7 (RPL7) mRNA, complete c
38 CATG GCTTTTAAGG A	-655115 L06498	Homo sapiens ribosomal protein S20 (RPS20) mRNA, c
GGCAA	-672265 L19527	Homo sapiens ribosomal protein L27 (RPL27) mRNA, c
	125346	- 1
40 CATG CTCTTCGAGA A	-490889 Y00433	Human mRNA for glutathione peroxidase (EC 1.11.1.9
	Y00483	Human gene for gluthathione peroxidase.
	X13710	H. sapiens unspliced mRNA for glutathione peroxidas
	X13709	Human gpx1 mRNA for gluthatione peroxidase.
	M21304	Human glutathione peroxidase (GPXI) mRNA, complete
41 CATG CTGTTGATTG C	-507455 X04347	Human liver mRNA fragment DNA binding protein UPI
	000947	Human clone C4E 3.2 (CAC)n/(GTG)n repeat-containin
42 CATG CTGGGTTAAT A	-502724 MB1757	H. sapiens S19 ribosomal protein mRNA, complete cds
CTGGTA	-239533 X17206	Human mRNA for LLRep3.
CATG GATG	-583573 X59357	Human mRNA for Epstein-Barr virus small RNAs (EBER
	121756	Homo sapiens acute myeloid leukemia associated pro
	017652	Human mRNA for HBp15/L22, complete cds.
	\$76343	AML1EAP (translocation breakpoint) [human, chro
45 CATG CCTTCGAGAT C	-390692 014970	Human ribosomal protein S5 mRNA, complete cds.
46 CATG CTCCTCACCT G	-482584 016811	Human Bak mRNA, complete cds.
	023765	
47 CATG TGTGTTGAGA G	-978825 X16869	r elongation factor 1-alpha (clone
	X16872	Human DNA for elongation factor 1-alpha (clone lam
	X03558	Human mRNA for elongation factor l alpha subunit (
	D17182	
	D17245	region Mbol cDNA,
	017259	region Mbol cDNA,
	D17276	Human HepG2 3' region MboI cDNA, clone hmd6a12m3.

			100 1753	
			M29548	Human elongation factor 1-alpha (EFIA) mRNA, parti
			141490	Homo sapiens oncogene PII-1 mRNA, complete cds.
			141498	Homo sapiens oncogene PTI-1 mRNA, complete cds.
8.8	ARICATE TIACCATATE	A -988	-988366 057846	Human ribosomal protein L39 mRNA, complete cds.
0 4	49 CATG GCCTGCTGGG	C -621	-621035 X71973	H.sapiens GPx-4 mRNA for phospholipid hydroperoxid
50	50 CATG CCTCGGAAAA	T -383	-383489 226876	H.sapiens gene for ribosomal protein L38.
1 5	ST CATG TACAAGAGGA	A -803	803369 X69391	H.sapiens mRNA for ribosomal protein L6.
1			803369 017554	Human mRNA for DNA-binding protein, TAXREB107, com
1		-803	-803369 S71022	neoplasm-related C140 product (human, thyroid carc
52	CATG AACGACCTCG	T -24	-24951 V00598	
-		-24	-24951 V00599	Human mRNA fragment encoding beta-tubulin. (from c
53	53 CATG CCCTGCCTTG	T -358	358783 X55110	ΨI
54	CATG CCCAGGGAGA	A -346	-346761 038846	Human stimulator of TAR RNA binding (SRB) mkNA, co
			D16933	77 (
2.5	55 CATG AGCACCTCCA	G -148	-148949 Z11692	H. sapiens mRNA for elongation factor 2.
2,6	56 CATG CGCCGGAACA	C -416	416261 X73974	H.sapiens HRPL4 mRNA.
			023660	Human mRNA for ribosomal protein, complete cds.
15	STICATE CTARABABA	A -458	-458753 M33680	TAPA-1 mR
, 5		989- 5	-686319 009510	Human glycyl-tRNA synthetase mRNA, complete cds.
3			009587	Human glycyl-tRNA synthetase mRNA, complete cds.
T			030658	Human T-cell mRNA for glycyl tRNA synthetase, comp
9,5	CATG ATTCTCCAGT	A -253	-253260 X55954	Human mRNA for HL23 ribosomal protein homologue.
;			X52839	Human mRNA for ribosomal protein L17.
60	ANICATE GAAAAATGGT	T -524	-524524 X61156	H.sapiens mRNA for laminin-binding protein.
3			X15005	Human mRNA for potential laminin-binding protein (
T			043901	Human 37 kD laminin receptor precursor/p40 ribosom
T			303799	Human colin carcinoma laminin-binding protein mRNA
			M14199	Ruman laminin receptor (2H5 epitope) mRNA, 5' end.
15	CATG CAGCTCACTG	A -302	-302367 D87735	Human mRNA for ribosomal protein L14, complete cds
			L10376	Human (clone CTG-B33) mRNA sequence.
			880520	CAG-is1 7 (trinucleotide repeat-containing sequenc
1	Entropy	200-	2005761114973	Human ribosomal protein S29 mRNA, complete cds.

					131610	Homo sapiens (clone cori-1cl5) \$29 ribosomal prote
12	CATG	63 CATG AATCCTGTGG	A	-55227	-55227 228407	H.sapiens mRNA for ribosomal protein L8.
9	CATG	64 CATG AATAGGTCCA	Æ	-51925	-51925 M64716	Human ribosomal protein S25 mRNA, complete cds.
			A (C,		-	
65	CATG	65 CATG AAAAAAAAA	G, T)	-1	-1 X83412	H.sapiens B1 mRNA for mucin.
					232564	H.sapiens FRGAMMA mRNA (819bp) for folate receptor
					232633	H.sapiens FRGAMMA' mRNA for folate receptor (817bp
					X76180	H.sapiens mRNA for lung amiloride sensitive Na+ ch
					U08470	Human FR-gamma' mRNA, complete cds.
					U08471	Human folate receptor 3 mRNA, complete cds.
					048697	Human mariner-like element-containing mRNA, clone
					D28532	Human mRNA for renal Na+-dependent phosphate cotra
					M55914	Human c-myc binding protein (MBP-1) mRNA, complete
					106175	Homo Sapiens P5-1 mRNA, complete cds.
					573775	calmitine=mitochondrial calcium-binding protein (h
					877393	transcript ch138 [human, RF1, RF48 stomach cancer c
					x60036	H.sapiens mRNA for mitochondrial phosphate carrier
200	CATG	SELCATG CCAGAACAGA	U	-335945 X79238	X79238	H.sapiens mRNA for ribosomal protein L30.
					Г16991	Human thymidylate kinase (CDC8) mRNA, complete cds
19	CATG	67 CATG AAGGTGGAGG	A	-44683 X80822	X80822	H.sapiens mRNA for ORF.
3 8	CATG	68 CATG CCTAGCTGGA	E	-379369 X52856	X52856	Human cyclophilin-related processed pseudogene.
					X52857	Human cyclophilin-related processed pseudogene.
					X52854	
					X52851	Human cyclophilin gene for cyclophilin (EC 5.2.1.8
					Y00052	Human mRNA for T-cell cyclophilin.
69	CATG	69 CATG GAACACATCC	A	-528694 X63527	X63527	H.sapiens mRNA for ribosomal protein L19.
					256985	ribosomal protein L19 [human, breast cancer cell l
10,5	CATG	70 CATG AAGGAGATGG	g	-41531 X69181	X69181	H.sapiens mRNA for ribosomal protein L31.
	_				X15940	Human mRNA for ribosomal protein L31.
7.1	71 CATG	AGGCTACGGA	ď.	-171113 229650	229650	H.sapiens SMCX mRNA.
					D17233	Human HepG2 3' region Mbol cDNA, clone hmd4c12m3.
72	CATG	72 CATG AGGTCCTAGC	U	-177610 X08096	96080X	Human GST pi gene for glutathione S-transferase pi
	_					

X15400 Human mRNN for anionic glutethione-S-transferase (Minds		X06547	Human mRNA for class Pi glutathione S-transferase
No. 10.287 Human glutathione S-transferase pi gene.		X15480	Human mRNA for anionic glutathione-S-transferase (
U12472 Human glutathione S-transferase (GST phi) gene, U21889 Human glutathione S-transferase Plo Gene, U05289 Human glutathione S-transferase Plo Gene, U05389 Human acidic ribosomal protein GHZ3] mkNA, U05389 Human acidic ribosomal protein GHZ3] mkNA, U05389 Human glutathione Gene, U05389 Human Glutathione Gene, U05389 Human Glutathione Gene, U05389 Human HegGZ 31 region Mboi Conglete Cds. U05480 Human Glutathione Gene, U05480 Human Gene, U05480		X08058	glutathione S-transferase pi
021689 Human glutathione S-transferase-Pic gene, compiled		012472	gene,
1062589 Human glutathione S-transferase Plo (GSTPLO) mit M69113 Human fatty acid ethyl ester synthase III mRNA M69113 Human fatty acid ethyl ester synthase III mRNA M6915 H. sapiens mRNA for ribosomal protein 51e.		021689	
Mig9113 Human fatty acid ethyl ester synthase-III mRNA Mig485 Homo sapiens (clone pHGST-pi) glutathione 5-tz Mig485 Homo sapiens spolipoprotein B gene sequence. Mig485 Homo sapiens apolipoprotein B gene sequence. Mig437 Homo sapiens 185 ribosomal protein (HKE31 mRNA L06437 Human acidic ribosomal protein (HKE31 mRNA L06438 Human acidic ribosomal protein (HKE31 mRNA L06439 Human acidic ribosomal protein PomENA, complete cds L06430 Human movel gene mRNA, complete cds L06431 Human movel gene mRNA, complete cds Mig431 Human movel gene mRNA, complete cds Mig431 Human acidic ribosomal protein (DN) mRNA, complete cds Mig431 Human movel gene mRNA, complete cds Mig431 Human acidic ribosomal protein (DN) mRNA, complete cds Mig440 Human mRNA for coupling protein G(s) alpha-sub Mig440 Human mRNA for coupling protein G-s, and color colo		062589	Human glutathione S-transferase Plc (GSTplc) mRNA,
W24485 Homo sapiens (clone pH6ST-pi) glutathione 5-tra W2465603 X65150 H.aapiens mRNA for ribosomal protein S18. W36153 Homo sapiens apolipoprotein B gene sequence. W36153 Homo sapiens 188 ribosomal protein (HKE3) mRNA W36154 Human acidic ribosomal protein PO mRNA, CATG GGGTTCCA -1769045 [25899 Human ribosomal protein LIO mRNA, complete cds W17371 Human W1855 DNA segment containing (TG)24 reg W17375 Human W1875 Human ribosomal protein LIO mRNA, complete cds W3756 Human W1875 Human W1875 Human, mEoceptor homolog (3' region) [Human, m		M69113	Human fatty acid ethyl ester synthase-III mRNA seg
CATG TGGTGTTGAG G -965603 X69150 H.sapiens mRNA for ribosomal protein S18.		M24485	glutathione
M96153 Homo sapiens apolipoprotein B gene sequence. 106422 Homo sapiens 185 ribosomal protein (HKE3) mRNA 106432 Homan acidic ribosomal protein (HKE3) mRNA, CAPIG GTGTTAACCA	CATG TGGTGTTGAG	-965603 X69150	
1.06432 Homeo sapiens 18S ribosomal protein (HKE3) mRNA		M96153	Homo sapiens apolipoprotein B gene sequence.
CATG GTCTAACATCT C -475448 M17885 Human acidic ribosomal phosphoprotein PO mRNA, CATG GTCTAACCA G -769045 L25899 Human ribosomal protein L10 mRNA, complete cds. CATG AGGCTTCCA A -174037 X58125 Human (D9S55) DNA segment containing (TG124 reg CATG AGGCTTCCA A -174037 X58125 Human (D9S55) DNA segment containing (TG124 reg CATG AGGCTTCCA A -174037 X58125 Human movel gene mRNA, complete cds. M13791 Human movel gene mRNA, complete cds. M17891 Human acidic ribosomal phosphoprotein P2 mRNA, CATG ATTAACAAAG C -671654 Human ferritin light subunit mRNA, partial cds. M11898 Human ferritin light subunit mRNA, complete cdc. M10119 Human ferritin light subunit mRNA, complete cdc. ATTAACAAAG C -246019 X04409 Human mRNA for coupling protein G(s) alpha-subunit AS5009 Human mRNA for alpha subunit of GsGTP binding protein alpha-m14631 Human guanine nucleotide-binding protein alpha-m2500 Human mRNA stimulatory GTP-binding protein G-s, alpha-m2500 Human mRNA stimulatory GTP-binding protein alpha-m2500 Human mRNA stimulatory GTP-binding protein alpha-m25000 Human mRNA stimulatory GTP-binding protein G-s, alpha-m2500 Human mRNA stimulatory GTP-binding protein G-s, alpha-tubulin mRNA, complete cds. M25009 Human M25000 Human M2-type pyruvate kinase mRNA, complete cds. Human M25000 Human M2-type pyruvate kinase mRNA, complete cds. Human M26252 Human TCB gene encoding cytosolic thyroid hormony M26252 Human TCB gene encoding cytosolic thyroid hormony M26200 Human TCB gene encoding cytosolic thyroid hormony M26200 Human TCB gene encoding cytosolic thyroid portein M26200 Human TCB gene encoding cytosolic thyroid portein M26200 Human TCB gene m2600 Human TCB gene m2600 Human TCB gene m2600 Human TCB gene M2600 Human M2600 Human TCB gene m2600 Human M2600 Human TCB gene m2600 Human Human Human Human TCB gene m2600 Human Human Human Human Human Human Human Human Human Human Human Hu		106432	
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Di7268 Human HepG2 3' region Mbol cDNA, clone hmd5h09m W13791 Human novel gene mRNA, complete cds. W64241 Human Wilm's tumnor-related protein (QM) mRNA, complete cds. S35960 laminin receptor homolog (3' region) [human, mb Human acidic ribosomal phosphoprotein P2 mRNA, complete cds. W11147 Human acidic ribosomal phosphoprotein P2 mRNA, complete cds. W11238 Human ferritin light subunit mRNA, complete cds. W1019 Human ferritin light subunit mRNA, complete cds. W1019 Human mRNA for coupling protein G(s) alpha-sub X5609 Human mRNA for coupling protein G(s) alpha-sub X5609 Human guanine nucleotide-binding protein alpha-luding protein G-s, a Human guanine nucleotide-binding protein G-s, a Human guanine nucleotide-binding protein G-s, a W14631 Human W2-type pyruvate kinase mRNA, complete cds W14631 Human M2-type pyruvate kinase mRNA, complete cd	CATG AGGGCTTCCA	-174037 X58125	Human (D9S55) DNA segment containing (TG)24 repeat
M13791 Human novel gene mRNA, complete cds. M64241 Human Wilm's tumor-related protein (QM) mRNA, complete cds. S35960 laminin receptor homolog {3' region} { human, mE } M1187 Human acidic ribosomal phosphoprotein P2 mRNA, mI M1187 Human ferritin L chain mRNA, complete cds. M12938 Human ferritin light subunit mRNA, complete cds. M10119 Human ferritin light subunit mRNA, complete cds. M10119 Human mRNA for coupling protein G(s) alpha-subuxof X36009 Human mRNA for coupling protein G(s) alpha-subuxof X36009 Human guanine nucleotide-binding protein alpha-m21142 Human guanine nucleotide-binding protein G-s, a M14631 Human guanine nucleotide-binding protein G-s, a M14631 Human alpha-tubulin mRNA, complete cds. K00558 Human alpha-tubulin mRNA, complete cds. K00558 Human alpha-tubulin mRNA, complete cds. M23725 Human M2-type pyruvate kinase mRNA, complete cds. M23725 Human M2-type pyruvate kinase mRNA, complete cds. M26552 Human TCB gene encoding cytosolic thyroid hormo		D17268	Human HepG2 3' region MboI cDNA, clone hmd5h09m3.
M64241 Human Wilm's tumor-related protein (QM) mRNA, CATG GGATTTGGCC T		M73791	Human novel gene mRNA, complete cds.
S35960 laminin receptor homolog (3' region) [human, mE		M64241	
CATG GGATTTGGCC F. 1654 M17887 Human acidic ribosomal phosphoprotein P2 mRNA, M11147 Human ferritin L chain mRNA, complete cds. CATG ATTAACAAAG C -246019 X04409 Human ferritin light subunit mRNA, complete cds CATG ATTAACAAAG C -246019 X04409 Human mRNA for coupling protein G(s) alpha-subunit xx56009 Human mRNA for coupling protein G(s) alpha-subunit xx56009 Human mRNA stimulatory GTP-binding protein alpha-subunit xx56009 M21142 Human guanine nucleotide-binding protein alpha-yx14631 M21142 Human guanine nucleotide-binding protein G-s, and yx14631 M14631 Human alpha-tubulin mRNA, 835bp. CATG TGGCCCCACC C -955718 X56494 H.sapiens M gene for MI-type and M2-type pyruvate kinase mRNA, complete cd M23725 Human M2-type pyruvate kinase mRNA, complete cd M26552 Human TCB gene encoding cytosolic thyroid hormonal ports		835960	(human,
M11147 Human ferritin L chain mRNA, complete cds.	CATG GGATTTGGCC	-671654 M17887	mRNA,
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CATG TGGCCCCACC C -955718 X56494 M23725 M26252	CATG TGTACCTGTA	-968173 236832	H.sapiens (xs31) mRNA, 835bp.
CATG TGGCCCCACC C -955718 X56494 M23725 M25252		K00558	human alpha-tubulin mRNA, complete cds.
M23725 M26252	CAT'G TGGCCCCACC	-955718 X56494	H.sapiens M gene for MI-type and M2-type pyruvate
		M23725	Human M2-type pyruvate kinase mRNA, complete cds.
		M26252	Human TCB gene encoding cytosolic thyroid hormone-

82	21.2		,			
	CATG		E	-602315 X89401	X89401	H.sapiens mRNA for large subunit of ribosomal prot
					014967	Human ribosomal protein L21 mRNA, complete cds.
	_				025789	Human ribosomal protein L21 mRNA, complete cds.
					L38826	Homo sapiens L21 ribosomal protein gene, partial c
7	CATG	TACCATCAAT	æ	-807748 X53778	X53778	H.sapiens hng mRNA for uracil DNA glycosylase.
					034995	Human normal keratinocyte substraction library mRN
	-				302642	Human glyceraldehyde 3-phosphate dehydrogenase mRN
					M36164	Human glyceraldehyde-3-phosphate dehydrogenase mRN
					M33197	Human glyceraldehyde-3-phosphate dehydrogenase (GA
8.4	CATG	ATTTGTCCCA	U	-260949 X14957	X14957	Human hmgI mRNA for high mobility group protein I.
;					X14958	Human hmgI mRNA for high mobility group protein Y.
					M23614	Human HMG-I protein isoform mRNA (HMGI gene), clon
					M23619	Human HMG-I protein isoform mRNA (HMGI gene), clon
					117131	Human high mobility group protein (HMG-I(Y)) gene
					M23615	Human HMG-Y protein isoform mRNA (HMGI gene), clon
					M23616	Human HMG-Y protein isoform mRNA (HMGI gene), clon
					M23617	Human HMG-Y protein isoform mRNA (HMGI gene), clon
					M23618	Human HMG-Y protein isoform mRNA (HMGI gene), clon
85	CATG	GAGGGAGTTT	υ	-567488 014968	U14968	Human ribosomal protein L27a mRNA, complete cds.
86	CATG	2992292292	ī	-416106012465	012465	Human ribosomal protein L35 mRNA, complete cds.
87	87 CATG GTGA	GTGAAACCCA	ALL	-753749 263072	263072	H.sapiens CpG island DNA genomic Msel fragment, cl
88	88 CATG GTGA	GTGAAACCCA	ALL	-753749 X16294	X16294	Human repetitive DNA containing interspersed repea
89	9 CATG	AAGACAGTGG	U	-33979	33979 X66699	- 1
					L06499	Homo sapiens ribosomal protein L37a (RFL37A) mRNA,
					L22154	Human ribosomal protein L37a mRNA sequence.
90	CATG	CCCCAGCCAG	t-	-348755 X55715	X55715	Human Hums3 mRNA for 40S ribosomal protein s3.
					014990	Human XP1PO ribosomal protein S3 (rpS3) mRNA, comp
	-				U14991	Human XP2NE ribosomal protein S3 (rpS3) mRNA, comp
			,		U14992	Human IMR-90 ribosomal protein S3 (rpS3) mRNA, com
	_				842658	S3 ribosomal protein [human, colon, mRNA, 826 nt].
ē	CATG	TGGGCAAAGC	U	-959498 X63526	X63526	H.sapiens mRNA for protein homologous to elongatio
	3				211531	H. sapiens mRNA for elongation factor-1-gamma.

92 CATG TGAGGGAATA A -928269 M10036 93 CATG GACGACACGA G -549145 U58682 94 CATG AACGCGGCCA A -26261 Z23063 95 CATG TGCACGTTTT C -935680 X03342 95 CATG TGCACGTTTT C -935680 X03342 96 CATG GGAGTGGACA T -667269 L119739 97 CATG GGGGAAATCG C -696375 M92381 100 CATG GCGCCATC G -696375 M92381 101 CATG TAAGGAGCTG A -796831 X77770 102 CATG GCCAACCCC A -796831 X77770 103 CATG GCCAACCCC A -6963742 U12474	
CATG TGAGGGAATA A CATG GACGACACGA G CATG AACGCGGCCA A CATG AACGCGGCCA A CATG CACAAACGGT A CATG GGAGTGGACA G CATG GGGGAAATCG C CATG GGGGAAACCCC A	Human triosephosphate isomerase mkNA, complete the them of the sound protein S28 mRNA, complete cds. Human ribosomal protein S4 (RPS4X) isoform mRNA Human scar protein mRNA, complete cds. Homo sapiens macrophage migration inhibitory ff Homo sapiens macrophage migration inhibitory ff Human migration inhibitory factor mRNA, complete cds Human mRNA for ribosomal protein L32. Human mRNA for ribosomal protein L32. Human ribosomal protein S27 mRNA, complete cds Homo sapiens metallopanstimulin (MPS1) mRNA, of Homo sapiens ribosomal protein L18 (RPL18) mRNA, complete CGS island DNA genomic Msel fragment H. sapiens CpG island DNA genomic Msel fragment H. sapiens CpG island DNA genomic Msel fragment H. sapiens CpG island DNA genomic Msel fragment H. sapiens CpG island DNA genomic Msel fragment H. sapiens CpG island DNA genomic Msel fragment H. sapiens CpG island DNA genomic Msel fragment H. sapiens CpG island DNA genomic Msel fragment H. sapiens CpG island DNA genomic Msel fragment
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CATG AACGCGGCCA A CATG TGCACGTTTT C CATG GCAAACGGT A CATG GCCAAACGGT G CATG GCCAAACGG C CATG GCCAAACCG C CATG GCCAAACCG C CATG GCCAAACCG C CATG GCCAAACCC A CATG TAAGGACCCC A CATG TAAGGACCCC A	Human ribosomal protein S4 (RPS4X) isoform mRNA, Human scar protein mRNA, complete cds. Homo sapiens macrophage migration inhibitory fact Homo sapiens macrophage migration inhibitory fact Homo sapiens macrophage migration inhibitory fact Human migration inhibitory fact Human mRNA for ribosomal protein L32. Human mRNA for ribosomal protein L32. Human mRNA from chromosome 15 gene with homology Human ribosomal protein S27 mRNA, complete cds. Homo sapiens metallopanstimulin (MPS1) mRNA, complement sapiens cpG island DNA genomic Msel fragment, of H. sapiens CpG island DNA genomic Msel fragment, of H. sapiens CpG island DNA genomic Msel fragment, of H. sapiens CpG island DNA genomic Msel fragment, of H. sapiens CpG island DNA genomic Msel fragment, of H. sapiens CpG island DNA genomic Msel fragment, of H. sapiens CpG island DNA genomic Msel fragment, of H. sapiens CpG island DNA genomic Msel fragment, of H. sapiens CpG island DNA genomic Msel fragment, of H. sapiens CpG island DNA genomic Msel fragment, or DNA ge
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CATG TGCACGTTTT C	Homo sapiens macrophage migration inhibitory fact Human migration inhibitory factor (MIF) mRNA, com Human mRNA for ribosomal protein L32. Human ribosomal protein S27 mRNA, complete cds. Homo sapiens metallopanstimulin (MPS1) mRNA, comp Homo sapiens ribosomal protein L18 (RPL18) mRNA, H. sapiens CpG island DNA genomic Msel fragment, c H. sapiens CpG island DNA genomic Msel fragment, c H. sapiens CpG island DNA genomic Msel fragment, c H. sapiens CpG island DNA genomic Msel fragment, c H. sapiens CpG island DNA genomic Msel fragment, c H. sapiens CpG island DNA genomic Msel fragment, c
CATG TGCACGTTTT C	Human migration inhibitory factor (MIF) mRNA, com Human mRNA for ribosomal protein L32. Human mRNA from chromosome 15 gene with homology Human ribosomal protein S27 mRNA, complete cds. Homo sapiens metallopanstimulin (MPS1) mRNA, comp Homo sapiens ribosomal protein L18 (RPL18) mRNA, H. sapiens CpG island DNA genomic Msel fragment, chromiens CpG island DNA genomic
CATG TGCACGTTTT C CATG CAAACGGT A CATG GGGGAAATGG C CATG GGGGAAATGG C CATG TAAGGAGCTG A CATG TAAGGAGCTG A CATG GGCAAGCCCC A CATG GGCAAGCCCC A CATG GGCAAGCCCC A CATG GGCAAGCCCC A CATG TGAAGCCCC A CATG TGAAGAACCCC A CATG TGAAGAACCCC A CATG TGAAGAACCCC A CATG TGAAGAACCCCC A CATG TGAAGAACCCC A CATG TGAAGAACCCC A CATG TGAAGAACCCC A CATG TGAAGAACCCCC A CATG TGAAGAACCCCC A CATG TGAAGAACCCC A CATG TGAAGAACCCCC A CATG TGAAGAACCCCC A CATG TGAAGAACCCCC A CATG TGAAGA	Human mRNA for ribosomal protein L32. Human mRNA from chromosome 15 gene with homology Human ribosomal protein S27 mRNA, complete cds. Homo sapiens metallopanstimulin (MPS1) mRNA, comp Homo sapiens ribosomal protein L18 (RPL18) mRNA, H.sapiens CpG island DNA genomic Msel fragment, c H.sapiens CpG island DNA genomic Msel fragment, c H.sapiens CpG island DNA genomic Msel fragment, c H.sapiens CpG island DNA genomic Msel fragment, c
CATG CACAAACGGT A CATG GGGGAAACG G CATG GGGGAAATCG C CATG GGGGAAATCG C CATG TAAGGAGCTG A CATG TAAGGAGCTC A	Human mRNA from chromosome 15 gene with homology Human ribosomal protein S27 mRNA, complete cds. Homo sapiens metallopanstimulin (MPS1) mRNA, comp Homo sapiens ribosomal protein L18 (RPL18) mRNA, H.sapiens CpG island DNA genomic Msel fragment, c H.sapiens CpG island DNA genomic Msel fragment, c H.sapiens CpG island DNA genomic Msel fragment, c H.sapiens CpG island DNA genomic Msel fragment, c
CATG CACAAACGGT A CATG GGAGTGGAAG G CATG GGGGAAATCG C CATG GGGGAAATCG C CATG TAAGGAGCTG A CATG TAAGGAGCCC A CATG GGCAAGCCCC A CATG GGCAAGCCCC A CATG GGCAAGCCCC A CATG CATG	Human ribosomal protein S27 mRNA, complete cds. Homo sapiens metallopanstimulin (MPS1) mRNA, comp Homo sapiens ribosomal protein L18 (RPL18) mRNA, H.sapiens CpG island DNA genomic Msel fragment, c H.sapiens CpG island DNA genomic Msel fragment, c H.sapiens CpG island DNA genomic Msel fragment, c
CATG GCAGTGGACA T CATG GCCGAGGAAG G CATG GGGGAAATCG C CATG GCAGCCATCC G CATG TAAGGAGCTG A CATG GCCAAGCCCC A	Homo sapiens metallopanstimulin (MPS1) mRNA, comp Homo sapiens ribosomal protein L18 (RPL18) mRNA, H.sapiens CpG island DNA genomic Msel fragment, o H.sapiens CpG island DNA genomic Msel fragment, o H.sapiens CpG island DNA genomic Msel fragment, o
CATG GCGAGGAAG G CATG GCCGAGGAAG G CATG GGGGAAATCG C CATG GCAGCCATCC G CATG TAAGGAGCTG A CATG GGCAAGCCCC A	Homo sapiens ribosomal protein L18 (RPL18) mRNA, H.sapiens CpG island DNA genomic Msel fragment, o H.sapiens CpG island DNA genomic Msel fragment, o H.sapiens CpG island DNA genomic Msel fragment, o
CATG GCGGAAGG G CATG GGGGAAATCG C CATG GCAGCCATC G CATG TAAGGAGCTG A CATG GGCAAGCCC A	H.sapiens CpG island DNA genomic Msel fragment, H.sapiens CpG island DNA genomic Msel fragment, H.sapiens CpG island DNA genomic Msel fragment,
CATG GGGGAAATCG C CATG GCAGCCATCC G CATG TAAGGAGCTG A CATG GGCAAGCCCC A	H.sapiens CpG island DNA genomic Msel fragment, H.sapiens CpG island DNA genomic Msel fragment,
CATG GGGGAAATCG C CATG GCAGCCATCC G CATG TAAGGAGCTG A CATG GGCAAGCCCC A	H. sapiens CpG island DNA genomic Msel fragment,
CATG GGGGAAATCG C CATG GCAGCCATCC G CATG TAAGGAGCTG A CATG GGCAAGCCCC A	2 - 2 - 2
CATG GGGGAAATCG C C CATG GCAGCCATCC G CATG TAAGGAGCTG A CATG GGCAAGCCCC A CATG GGCAAGCCCC A	X53505 Human mknA for fibusomai process sit:
CATG GCAGCCATCC G	92381 Human thymosin beta 10 mRNA, complete cds.
CATG GCAGCCATCC G — CATG TAAGGAGCTG A — CATG GGCAAGCCCC A —	M20259 Human thymosin beta-10 mRNA, complete cds.
CATG TAAGGAGCTG A -796831 CATG GGCAAGCCCC A -672342	14969 Human ribosomal protein L28 mRNA, complete cds.
CATG TAAGGAGCTG A -796831 CATG GGCAAGCCCC A -672342	D17257 Human HepG2 3' region Mbol cDNA, clone hmd5d04m3.
CATG GGCAAGCCCC A -672342	H.sapiens RPS26 mRNA.
CATG GGCAAGCCCC A -672342	X69654 H.sapiens mRNA for ribosomal protein S26.
X75	. S13.
101	L01124 Human ribosomal protein S13 (RPS13) mRNA, complete
103 CATG GTTCCCTGGC C -775658 X65923	
104 CATG CCGTCCAAGG G -374027 M60	M60854 Human ribosomal protein S16 mRNA, complete cds.
CATG TTGG	H. sapiens mRNA for homologue to yeast ribo
	S64030 [L41 ribosomal protein homolog (clone 786) [human,

105 CATG CAAACCATCC A	-263478 X12883	Human mRNA for cytokeratin 18.
	X12876	Human mRNA fragment for cytokeratin 18.
	X12881	Human mRNA for cytokeratin 18.
	M24842	Human keratin 18 (K18) gene, complete cds.
	M26325	Human cytokeratin 18 mRNA, 3' end.
	M26326	Human keratin 18 mRNA, complete cds.
	M26327	Human cytokeratin 18 mRNA, 3' end.
106 CATG AGCTCTCCCT G	-161624 X53777	Human 123 mRNA for putative ribosomal protein.
107 CATG AGGTCAGGAG A(T)	-177315 D86979	Human male bone marrow myeloblast mRNA for KIAA022
	X55923	Human DNA for Alu element PIN6.
	X79699	H.sapiens ALU repeat, 230bp.
	X12544	Human mRNA for HLA class II DR-beta (HLA-DR B).
	686112	H.sapiens flow~sorted chromosome 6 HindIII fragmen
	011831	Human clone 2102V-I chromosome 18p telomeric seque
	012580	Human Alu repeat sequence A3.
	012582	Human Alu repeat sequence B2.
	012583	Human Alu repeat sequence DI.
	U14694	Human Alu-Sb2 repeat, clone HALUSBOB.
	014695	Human Alu-Sb2 repeat, clone HALUSB15.
	014696	Human Alu-Sb2 repeat, clone HALUSB27.
	014697	Human Alu-Sb2 repeat, clone HUM-11.
	014698	Human Alu-Sb2 repeat, clone HSB-8P.
	014699	Human Alu-Sb2 repeat, clone HUM-9.
	014700	Human Alu-Sb2 repeat, clone HALUSB35.
	014701	- 1
	014704	Alu-Sb2 repeat, clone
	014706	Ruman Alu-Sb2 repeat, clone HUM-10.
	014707	Human Alu-Sb2 repeat, clone HUM-7.
	500120	Human (Lawn) c-myc proto-oncogene, complete coding
	134653	Homo sapiens platelet/endothelial cell adhesion mo
,	M37521	Human XV2c gene.
	861789	NFI-neurofibromatosis type 1 (deletion breakpoint,
	873483	phosphorylase kinase catalytic subunit PHKG2 homol

				875201	cholinesterase (Alu element) [human, Insertion Mut
				875337	(Y Alu polymorphism, YAP, polymorphic subfamily-3}
108	108 CATG GGGCTGGGGT	O J	-695980	695980 249148	H.sapiens mRNA for ribosomal protein 129.
				U10248	Human ribosomal protein L29 (humrp129) mRNA, compl
				049083	Human cell surface heparin binding protein HIP mRN
				D16992	Human HepG2 partial cDNA, clone hmd2d02m5.
				D16911	Human HepG2 3' region cDNA, clone hmd3b09.
				J03537	Human ribosomal protein S6 mRNA, complete cds.
				M20020	Ruman ribosomal protein S6 mRNA, complete cds.
109	109 CATG ACGTTCTCTT	O	-114144		EST
110	110 CATG TCTCCATACC	U	-906438		EST
111	111 CATG GACTGCGTGC	O	-555450		EST
112	112 CATG CTTAATCCTG	ď	-508767		EST
113	113 CATG GGTTGGCAGG	9	-719435		EST
114	114 CATG GCCCTCTGCC	A	-613862		EST
115	115 CATG AACAGAAGCA	e 1	-18469		EST
116	116 CATG CTGCCGAGCT	υ	-497192		EST
1117	117 CATG TTCCTCGGGC	A	-1007018		EST
118	118 CATG AACTAATACT	A	-28872		EST
119	119 CATG TAGATAATGG	υ	-822331		EST
120	120 CATG GCCACACCCC	A,C	-607318		EST
121	121 CATG GAACCCTGGG	A	-529899		EST
122	122 CATG AACTAAAAA	A A	-28673		EST
123	123 CATG GAAATGTAAG	Ą	-528067		EST
124	124 CATG ACTCCAAAAA	A 1	-119809		EST
125	125 CATG GTTCGTGCCA	A	-777109		EST
126	126 CATG TTACCTCCTT	٠ ,	-989024		EST
127	127 CATG GCACAAGAAG	A	-594051	-	EST
128	128 CATG CCCTGGGTTC	T	-359102		EST
129	129 CATG GCCTGTATGA	9	-621369	`	EST
130	130 CATG CCCGTCCGGA	A	-355689		EST
131	131 CATG AGGAAAGCTG	<u>ن</u>	~163999		EST
132	132 CATG TCAGATCTTT	υ	-861056		EST

EST EST EST EST

			۱	-	1338081
_	133	123 CATG	CCAGGAGGAA T		10000
	1			-	-857362
-	1 2/	124 6476	TCACCACAC		
	1			-	-769605
-	1.00	TACIDAT	A WORDSTITED A		
_	7	2		-	618195
_	36	136 CATG	GCCGTGTCCG		2070
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Isolation of partial cDNA (3' fragment) by 3' directed PCR reaction

This procedure is a modification of the protocol described in Polyak et al. (1997) Nature 389:300. Briefly, the procedure uses SAGE tags in PCR reaction such that the resultant PCR product contains the SAGE tag of interest as well as additional cDNA, the length of which is defined by the position of the tag with respect to the 3' end of the cDNA. The cDNA product derived from such a transcript driven PCR reaction can be used for many applications.

RNA from a source believed to express the cDNA corresponding to a given tag is first converted to double-stranded cDNA using any standard cDNA protocol. Similar conditions used to generate cDNA for SAGE library construction can be employed except that a modified oligo-dT primer is used to dreive the first strand synthesis. For example, the oligonucleotide of compositon 5'-B-TCC GGC GCG CCG TTT T CC CAG TCA CGA(30)-3', contains a poly-T stretch at the 3' end for hybridization and priming from poly-A tails, an M13 priming site for use in subsequent PCR steps, a 5' Biotin label (B) for capture to strepavidin-coated magnetic beads, and an AscI restriction endonuclease site for releasing the cDNA from the streptavidin-coated magnetic beads. Theoretically, any sufficiently-sized DNA region capable of hybridizing to a PCR primer can be used as well as any other 8 base pair recognizing endonuclease.

cDNA constructed utilizing this or similar modified oligo-dT primer is then processed exactly as described in U.S. Patent No. (insert) up until adapter ligation where only one adapter is ligated to the cDNA pool. After adapter ligation, the cDNA is released from the streptavidin-coated magnetic beads and is then used as a template for cDNA amplification.

Various PCR protocols can be employed using PCR priming sites within the 3' modified oligo-dT primer and the SAGE tag. The SAGE tag-derived PCR primer employed can be of varying length dictated by 5' extension of the tag into the adaptor sequence. cDNA products are now available for a variety of applications.

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This technique can be further modified by: (1) altering the length and/or content of the modified oligo-dT primer; (2) ligating adaptors other than that previously employed within the SAGE protocol; (3) performing PCR from template retained on the streptavidin-coated magnetic beads; and (4) priming first strand cDNA synthesis with non-oligo-dT based primers.

Isolation of cDNA using GeneTrapper or modified GeneTrapper Technology

The reagents and manufacturer's instructions for this technology are commercially available from Life Technologies, Inc., Gaithersburg, Maryland. Briefly, a complex population of single-stranded phagemid DNA containing directional cDNA inserts is enriched for the target sequence by hybridization in solution to a biotinylated oligonucleotide probe complementary to the target sequence. The hybrids are captured on streptavidin-coated paramagnetic beads. A magnet retrieves the paramagnetic beads from the solution, leaving nonhybridized single-stranded DNAs behind. Subsequently, the captured single-stranded DNA target is released from the biotinylated oligonucleotide. After release, the cDNA clone is further enriched by using a nonbiotinylated target oligonucleotide to specifically prime conversion of the single-stranded target to double-stranded DNA. Following transformation and plating, typically 20% to 100% of the colonies represent the cDNA clone of interest. To identify the desired cDNA clone, the colonies may be screened by colony hybridization using the 32P-labeled oligonucleotide as described above for solution hybridization, or alternatively by DNA sequencing and alignment of all sequences obtained from numerous clones to determine a consensus sequence.

The genes which are identified herein as being differentially expressed in normal and cancer cells can be used diagnostically and prognostically. Transcription levels in a test sample suspected of being neoplastic can be determined and compared to the levels in normal colon cells. The test sample may be from any tissue suspected of neoplasia, and particularly from either suspected colorectal or suspected pancreatic cancer cells. The control cells for

the purposes of comparison are normal cells, preferably of the same tissue type as the test sample, e.g., colon cells, or pancreatic duct epithelial cells. Upregulation of transcription or downregulation of transcription is therefore diagnostic of the neoplastic state, depending on what gene is used as a test reagent. Similarly, transcription levels can be monitored to assess patent responses to anti-tumor therapies. Transcription levels will also provide prognostic information. For example, the level of transcription in a test sample can be compared to levels found in bona fide normal and tumor cells. More extreme deviations from normal expression levels indicate a poorer prognosis.

Transcription levels can be determined according to any means known in the art. These include, without limitation, Northern blots, nuclear run-on assays, *in vitro* transcription assays, primer extension assays, quantitative reverse transcriptase-polymerase chain reactions (RT-PCR), and hybrid filter binding assays. These techniques are well known in the art. See J.C. Alwine, D.J. Kemp, G.R. Stark, *Proc. Natl. Acad. Sci. U.S.A.* 74, 5350 (1977); K. Zinn, D. Di-Maio, T. Maniatis, *Cell* 34, 865 (1983); G. Veres, R.A. Gibbbs, S.E. Scherer, C.T. Caskey, *Science* 237, 415 (1987).

Similarly, upregulated genes and downregulated genes can be detected by measuring expression of their protein products. This can be done by any means known in the art, including but not limited to Western (immuno) blot, enzyme linked immunoadsorbent assay, radioimmunoassay, and enzyme assay. Such techniques are well known in the art. Protein products can be detected in tissue samples of a test patient, using a suspect sample as a test sample, and a matched normal tissue sample from the same tissue type as a control. If normal tissue is not available then a closely related tissue type can be used. Desirably both the samples being compared will be from the same individual. Alternatively, aberrant expression levels of protein products can be detected in body samples, such as blood, serum, feces, urine, sputum. As a control, a normal matched sample can be used from a healthy individual. Aberrant expression levels of transcripts can also be detected in such body samples, particularly in blood and serum.

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Probes for use in the assays for transcription levels of particular genes or sets of genes may be RNA or DNA. The probes will be isolated substantially free of other cellular RNAs or DNAs. If the reagent contains one probe then it will comprise at least 50% of the nucleic acids in the reagent composition. If the reagent contains more than one probe, then the proportion will decrease accordingly, so that specific probes will still comprise at least 50% of the nucleic acids in the reagent composition.

Probes can be labeled according to any means known in the art. These may include radioactive labels, fluorescent labels, enzymatic labels, and binding partner labels such as biotin. Means for labeling and detecting probes are well known in the art. Probes comprise at least 10, 11, 12, 15, 20, or 30 contiguous nucleotides of a selected gene.

This invention provides proteins or polypeptides expressed from the polynucleotides of this invention, which is intended to include wild-type and recombinantly produced polypeptides and proteins from procaryotic and eucaryotic host cells, as well as muteins, analogs and fragments thereof. In some embodiments, the term also includes antibodies and anti-idiotypic antibodies.

It is understood that functional equivalents or variants of the wild-type polypeptide or protein also are within the scope of this invention, for example, those having conservative amino acid substitutions. Other analogs include fusion proteins comprising a protein or polypeptide.

The proteins and polypeptides of this invention are obtainable by a number of processes well known to those of skill in the art, which include purification, chemical synthesis and recombinant methods. Full length proteins can be purified from a colon or pancreatic cell or tissue lysate by methods such as immunoprecipitation with antibody, and standard techniques such as gel filtration, ion-exchange, reversed-phase, and affinity chromatography using a fusion protein as shown herein. For such methodology, see for example Deutscher et al. (1999) Guide To Protein Purification: Methods In Enzymology (Vol. 182, Academic Press). Accordingly, this invention also

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provides the processes for obtaining these proteins and polypeptides as well as the products obtainable and obtained by these processes.

The proteins and polypeptides also can be obtained by chemical synthesis using a commercially available automated peptide synthesizer such as those manufactured by Perkin Elmer/Applied Biosystems, Inc., Model 430A or 431A, Foster City. The synthesized protein or polypeptide can be precipitated and further purified, for example by high performance liquid chromatography (HPLC). Accordingly, this invention also provides a process for chemically synthesizing the proteins of this invention by providing the sequence of the protein and reagents, such as amino acids and enzymes and linking together the amino acids in the proper orientation and linear sequence.

Alternatively, the proteins and polypeptides can be obtained by well-known recombinant methods as described, for example, in Sambrook et al., (1989), supra, using the host cell and vector systems described above.

Also provided by this application are the polypeptides and proteins described herein conjugated to a detectable agent for use in the diagnostic methods. For example, detectably labeled proteins and polypeptides can be bound to a column and used for the detection and purification of antibodies. They also are useful as immunogens for the production of antibodies as described below. The proteins and fragments of this invention are useful in an in vitro assay system to screen for agents or drugs, which modulate cellular processes.

The proteins of this invention also can be combined with various liquid phase carriers, such as sterile or aqueous solutions, pharmaceutically acceptable carriers, suspensions and emulsions. Examples of non-aqueous solvents include propyl ethylene glycol, polyethylene glycol and vegetable oils. When used to prepare antibodies, the carriers also can include an adjuvant that is useful to non-specifically augment a specific immune response. A skilled artisan can easily determine whether an adjuvant is required and select one. However, for the purpose of illustration only, suitable adjuvants include, but

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are not limited to Freund's Complete and Incomplete, mineral salts and polynucleotides.

This invention also provides a pharmaceutical composition comprising any of a protein, analog, mutein, polypeptide fragment, antibody, antibody fragment or anti-idiotipic antibody of this invention, alone or in combination with each other or other agents, and an acceptable carrier. These compositions are useful for various diagnostic and therapeutic methods.

Antibodies can be generated using the proteins encoded by the transcripts identified by the tags disclosed herein. Use of all or portions of the protein as immunogens is routine in the art. Similarly, fusion proteins can be used as immunogens. Antibodies can be affinity purified using the proteins or portions thereof used as immunogens. Similarly, monoclonal antibodies specifically immunoreactive with the protein sequences of the invention can be generated according to techniques which are well known in the art.

Antibodies can be used analytically to quantitate the expression of particular transcripts identified herein as upregulated or downregulated in cancer. In addition, antibodies can be conjugated or non-covalently linked to cytotoxic agents, such as cytotoxins, radionuclides, chemotherapeutic drugs, etc. Such antibodies can be used therapeutically to specifically target cancer cells in which the protein antigens are upregulated. These include the proteins encoded by the transcripts identified by the tags shown in Tables 2, 4, and 5. Means of making such linked cytotoxic antibodies and of administering the same are well known in the art.

Also provided by this invention is an antibody capable of specifically forming a complex with the proteins or polypeptides as described above. The term "antibody" includes polyclonal antibodies and monoclonal antibodies. The antibodies include, but are not limited to mouse, rat, and rabbit or human antibodies.

Laboratory methods for producing polyclonal antibodies and monoclonal antibodies, as well as deducing their corresponding nucleic acid sequences, are known in the art, see Harlow and Lane (1988) supra and

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Sambrook et al. (1989) supra. The monoclonal antibodies of this invention can be biologically produced by introducing protein or a fragment thereof into an animal, e.g., a mouse or a rabbit. The antibody producing cells in the animal are isolated and fused with myeloma cells or heteromyeloma cells to produce hybrid cells or hybridomas. Accordingly, the hybridoma cells producing the monoclonal antibodies of this invention also are provided.

Thus, using the protein or fragment thereof, and well known methods, one of skill in the art can produce and screen the hybridoma cells and antibodies of this invention for antibodies having the ability to bind the proteins or polypeptides.

If a monoclonal antibody being tested binds with the protein or polypeptide, then the antibody being tested and the antibodies provided by the hybridomas of this invention are equivalent. It also is possible to determine without undue experimentation, whether an antibody has the same specificity as the monoclonal antibody of this invention by determining whether the antibody being tested prevents a monoclonal antibody of this invention from binding the protein or polypeptide with which the monoclonal antibody is normally reactive. If the antibody being tested competes with the monoclonal antibody of the invention as shown by a decrease in binding by the monoclonal antibody of this invention, then it is likely that the two antibodies bind to the same or a closely related epitope. Alternatively, one can pre-incubate the monoclonal antibody of this invention with a protein with which it is normally reactive, and determine if the monoclonal antibody being tested is inhibited in its ability to bind the antigen. If the monoclonal antibody being tested is inhibited then, in all likelihood, it has the same, or a closely related, epitopic specificity as the monoclonal antibody of this invention.

The term "antibody" also is intended to include antibodies of all isotypes. Particular isotypes of a monoclonal antibody can be prepared either directly by selecting from the initial fusion, or prepared secondarily, from a parental hybridoma secreting a monoclonal antibody of different isotype by using the sib selection technique to isolate class switch variants using the

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procedure described in Steplewski et al. (1985) Proc. Natl. Acad. Sci. 82:8653 or Spira et al. (1984) J. Immunol. Methods 74:307.

This invention also provides biological active fragments of the polyclonal and monoclonal antibodies described above. These "antibody fragments" retain some ability to selectively bind with its antigen or immunogen. Such antibody fragments can include, but are not limited to:

- (1) Fab,
- (2) Fab',
- (3) F(ab')2,
- (4) Fv, and
- (5) SCA

A specific example of "a biologically active antibody fragment" is a CDR region of the antibody. Methods of making these fragments are known in the art, see for example, Harlow and Lane, (1988) supra.

The antibodies of this invention also can be modified to create chimeric antibodies and humanized antibodies (Oi, et al. (1986) BioTechniques 4(3):214). Chimeric antibodies are those in which the various domains of the antibodies' heavy and light chains are coded for by DNA from more than one species.

The isolation of other hybridomas secreting monoclonal antibodies with the specificity of the monoclonal antibodies of the invention can also be accomplished by one of ordinary skill in the art by producing anti-idiotypic antibodies (Herlyn, et al. (1986) Science 232:100). An anti-idiotypic antibody is an antibody which recognizes unique determinants present on the monoclonal antibody produced by the hybridoma of interest.

Idiotypic identity between monoclonal antibodies of two hybridomas demonstrates that the two monoclonal antibodies are the same with respect to their recognition of the same epitopic determinant. Thus, by using antibodies to the epitopic determinants on a monoclonal antibody it is possible to identify other hybridomas expressing monoclonal antibodies of the same epitopic specificity.

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It is also possible to use the anti-idiotype technology to produce monoclonal antibodies which mimic an epitope. For example, an anti-idiotypic monoclonal antibody made to a first monoclonal antibody will have a binding domain in the hypervariable region which is the mirror image of the epitope bound by the first monoclonal antibody. Thus, in this instance, the anti-idiotypic monoclonal antibody could be used for immunization for production of these antibodies.

As used in this invention, the term "epitope" is meant to include any determinant having specific affinity for the monoclonal antibodies of the invention. Epitopic determinants usually consist of chemically active surface groupings of molecules such as amino acids or sugar side chains and usually have specific three dimensional structural characteristics, as well as specific charge characteristics.

The antibodies of this invention can be linked to a detectable agent or label. There are many different labels and methods of labeling known to those of ordinary skill in the art.

The antibody-label complex is useful to detect the protein or fragments in a sample, using standard immunochemical techniques such as immunohistochemistry as described by Harlow and Lane (1988) supra. Competitive and non-competitive immunoassays in either a direct or indirect format are examples of such assays, e.g., enzyme linked immunoassay (ELISA) radioimmunoassay (RIA) and the sandwich (immunometric) assay. Those of skill in the art will know, or can readily discern, other immunoassay formats without undue experimentation.

The coupling of antibodies to low molecular weight haptens can increase the sensitivity of the assay. The haptens can then be specifically detected by means of a second reaction. For example, it is common to use haptens such as biotin, which reacts avidin, or dinitropherryl, pyridoxal, and fluorescein, which can react with specific anti-hapten antibodies. See Harlow and Lane (1988) supra.

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The monoclonal antibodies of the invention also can be bound to many different carriers. Thus, this invention also provides compositions containing the antibodies and another substance, active or inert. Examples of well-known carriers include glass, polystyrene, polypropylene, polyethylene, dextran, nylon, amylases, natural and modified celluloses, polyacrylamides, agaroses and magnetite. The nature of the carrier can be either soluble or insoluble for purposes of the invention. Those skilled in the art will know of other suitable carriers for binding monoclonal antibodies, or will be able to ascertain such, using routine experimentation.

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Compositions containing the antibodies, fragments thereof or cell lines which produce the antibodies, are encompassed by this invention. When these compositions are to be used pharmaceutically, they are combined with a pharmaceutically acceptable carrier.

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The present invention also provides a screen for various agents which modulate the expression of a gene in a pancreatic or colon cell. To practice the method in vitro, suitable cell cultures or tissue cultures are first provided. The cell can be a cultured cell or a genetically modified cell in which a trancript from SEQ ID NOS:1-732, or their complements, is expressed. Alternatively, the cells can be from a tissue biopsy. The cells are cultured under conditions (temperature, growth or culture medium and gas (CO₂)) and for an appropriate amount of time to attain exponential proliferation without density dependent constraints. It also is desirable to maintain an additional separate cell culture; one which does not receive the agent being tested as a control.

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As is apparent to one of skill in the art, suitable cells may be cultured in microtiter plates and several agents may be assayed at the same time by noting genotypic changes, phenotypic changes or cell death.

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When the agent is a composition other than a DNA or RNA, the agent may be directly added to the cell culture or added to culture medium for addition. As is apparent to those skilled in the art, an "effective" amount must be added which can be empirically determined. When the agent is a polynucleotide, it may be directly added by use of a gene gun or

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electroporation. Alternatively, it may be inserted into the cell using a gene delivery vehicle or vector as described above.

An agent is a potential therapeutic if it alters the expression of gene in the cell. Altered expression can be detected by assaying for altered mRNA expression or protein expression using the probes, primers and antibodies as described herein.

For the purposes of this invention, an "agent" is intended to include, but not be limited to a biological or chemical compound such as a simple or complex organic or inorganic molecule, a peptide, a protein (e.g. antibody) or a polynucleotide (e.g. anti-sense). A vast array of compounds can be synthesized, for example polymers, such as polypeptides and polynucleotides, and synthetic organic compounds based on various core structures, and these are also included in the term "agent". In addition, various natural sources can provide compounds for screening, such as plant or animal extracts, and the like. It should be understood, although not always explicitly stated that the agent is used alone or in combination with another agent, having the same or different biological activity as the agents identified by the inventive screen. The agents and methods also are intended to be combined with other therapies.

The above disclosure generally describes the present invention. A more complete understanding can be obtained by reference to the following specific examples which are provided herein for purposes of illustration only, and are not intended to limit the scope of the invention.

EXAMPLE 1

This example demonstrates the characterization of the general transcription of human colorectal epithelium, colorectal cancers, and pancreatic cancers.

We used the recently developed SAGE (serial analysis of gene expression) method to identify and quantify a total of 303,706 transcripts derived from human colorectal (CR) epithelium, CR cancers or pancreatic cancers (Table 1A) (3). These transcripts represented approximately 48,741

different genes (4) that ranged in average expression from 1 copy per cell to as many as 5,300 copies per cell (5). The number of different transcripts observed in each cell population varied from 14,247 to 20,471. The bulk of the mRNA mass (75%) consisted of transcripts expressed at more than five copies per cell on average (Table 1B). In contrast, the majority (86%) of transcripts were expressed at less than 5 copies per cell, but in aggregate this low abundance class represented only 25% of the mRNA mass. This distribution was consistently observed among the different samples analyzed and was consistent with previous studies of RNA abundance classes based on RNA-DNA reassociation kinetics (Rot curves). Monte Carlo simulations revealed that our analyses had a 92% probability of detecting a transcript expressed at an average of three copies per cell (7).

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Table 1 - Summary of SAGE Analysis

A. Overall Summary

	Normal	Colon	Colon	Pancreatic	Pancreatic	
	Colon	Tumors	Cell Lines	Tumors	Cell Lines	Total
Total Tags	62,168	80,878	60,373	61,592	58,695	303,706
Unique Genes¹ GenBank²	14,721 8,753 (59)	19,690 10,490 (53)	17,092 10,193 (60)	20,471 11,547 (56)	14,247 8,922 (63)	48,741 26,339 (54)

¹ Indicates the number of different genes represented by the total tags analyzed (4).

² Indicates the number of genes that matched an entry in GenBank. The number in parentheses indicates the corresponding percentage of total unique tags.

Table 1 - Summary of SAGE Analysis

B. Summarized by Abundance Classes*

	Normal	Colon	Colon	Pancreatic	Pancreatic Cell	
Copies/Cell	Colon	Tumors	Cell Lines	Tumors	Lines	Total
> 500						
Unique Genes	62 (29)	54 (25)	54 (19)	32 (11)	70 (26)	55 (19)
GenBank	(56) 65	52 (96)	53 (98)	32 (100)	70 (100)	54 (98)
> 50 and < 500		į	Í	ļ	ĺ	; ;
Unique Genes	645 (28)	470 (21)	618 (27)	657 (29)	585 (27)	595 (26)
GenBank	545 (84)	429 (91)	579 (94)	(66) (63)	529 (90)	553 (93)
> 5 and < 50						
Unique Genes	4,569 (27)	5,011 (29)	5,733 (34)	6,146 (36)	4,895 (31)	6,209 (30)
GenBank	2,893 (63)	3,204 (64)	3,682 (64)	4,054 (66)	3,168 (65)	4,241 (68)

41,882 (25)	21,491 (51)
8,697 (16)	5,155 (59)
13,636 (24)	6,852 (50)
10,687 (20)	5,879 (55)
14,155 (25)	6,805 (48)
9,445 (16)	5,256 (56)
< 5 Unique Genes	GenBank

*For unique genes, the first number denotes the number of different genes (4) represented in the indicated abundance class. The number in parentheses indicates the mass fraction (X100) of total transcripts represented by the indicated abundance class. For GenBank entries, the first number indicates the number of different genes that matched an entry in GenBank in the indicated abundance class. The number in parentheses indicates the corresponding percentage of total genes.

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Many of the SAGE tags appeared to represent previously undescribed transcripts, as only 54% of the tags matched entries in GenBank (Table 1). Twenty percent of these matching transcripts corresponded to characterized mRNA sequence entries in GenBank, whereas 80% matched uncharacterized EST entries. As expected, the likelihood of a tag being present in the databases was related to abundance; GenBank matches were identified for 98% of the transcripts expressed at more than 500 copies per cell but for only 51% of the transcripts expressed at \leq 5 copies per cell. Because the SAGE data provide a quantitative assay of transcript abundance, unaffected by differences in cloning or PCR efficiency, these data provide an independent and relatively unbiased estimate of the current completeness of publicly available EST databases.

EXAMPLE 2

This example demonstrates a comparison of the expression pattern of normal colon epithelium and primary colon cancers.

Comparison of expression patterns between normal colon epithelium and primary colon cancers revealed that the majority of transcripts were expressed at similar levels (Fig. 1A). However, the expression profiles also revealed 289 transcripts that were expressed at significantly different levels [P < 0.01, (8)]. Of these 289, 181 were decreased in colon tumors compared to normal colon (average decrease 10-fold; Fig. 1B; examples in Fig. 2A). Conversely, 108 transcripts were expressed at higher levels in the colon cancers than in normal colon (average increase 13-fold; Fig. 1C; examples in Fig. 2A). Monte Carlo simulations indicated that the analysis would have detected over 95% of those transcripts expressed at a 6-fold or greater level in normal vs. tumor cells or vice versa (9). Because relatively stringent criteria were used for defining differences [P < 0.01, (8)], the number of differences reported above is likely to be an underestimate.

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EXAMPLE 3

This example demonstrates the similarities and differences between cancer cell line transcription and transcription of primary cancer tissues. To determine how many of the 289 differences were independent of the cellular microenvironment of cancers in vivo, SAGE data from CR cancer cell lines was compared to that from primary CR cancer tissues (Fig. 1B, 1C). Perhaps surprisingly, the majority of transcripts (130 of 181) that were expressed at reduced levels in cancer cells in vivo were also expressed at significantly lower levels in the cell lines (Fig. 1B). Likewise, a significant fraction of the transcripts expressed at increased levels in primary cancers were also expressed at higher levels in the CR cancer cell lines (Fig. 1C). Thus, many of the gene expression differences that distinguish normal from tumor cells in vivo persist during in vitro growth. However, despite these similarities there were also many differences. For example, only 47 of 228 genes expressed at higher levels in CR cancer cell lines were also expressed at high levels in the primary CR cancers.

In combination, comparing the expression pattern of CR cancer cells (in vivo or in vitro) to normal colon revealed 548 differentially expressed transcripts (Fig. 1B,C, Tables 2 and 3). The average difference in expression for these transcripts was 15 fold. Although the ability to detect differences is influenced by the magnitude of the variance with the power to detect smaller differences being less, 92 transcripts that were less than three fold different were identified among the 548 transcripts. However, those genes exhibiting the greatest differences in expression are likely to be the most biologically important.

EXAMPLE 4

This example demonstrates the similarities and differences between colorectal cancer transcription and pancreatic cancer transcription.

To determine whether the changes noted in CR cancers were neoplasia or cell type specific, we performed SAGE on mRNA derived from pancreatic cancers. A total of 404 transcripts were expressed at higher levels in pancreatic cancers compared to normal colon epithelium (examples in Fig. 2B). The majority (268) of these transcripts were pancreas-specific (10) (Example in Fig. 2C) although 136 were also expressed at high levels in CR cancers. These 136 transcripts constituted 47% of the 289 transcripts increased in CR cancers relative to normal colon and are likely to be related to the neoplastic process rather than to the specific cell type of origin.

EXAMPLE 5

This example demonstrates the reproducibility of the transcription patterns observed among a larger number of cancer samples.

One question that arose from these data is the potential heterogeneity of expression between individual tumors. The SAGE data were acquired from two examples of each tissue type (normal colon, primary CR cancer, CR cancer cell line, etc.). To examine the generality of these expression profiles, we arbitrarily selected 27 differentially expressed transcripts and evaluated them in six to twelve samples of normal colon and primary cancers by Northern blot analysis (11). In general, expression patterns were very reproducible among different samples. Of 10 genes with elevated expression in normal colon relative to CR cancers as determined by SAGE, each was detected in the normal colon samples and was expressed at considerably lower levels in tumors (examples in Fig. 2A). Similarly, most of the genes identified by SAGE as increased in CR or pancreatic cancers were confirmed to be reproducibly expressed in the majority of primary cancers examined by Northern blot (examples in Fig. 2A). It is important to note, however, that there were differences among the cancers, with a few cancers exhibiting particularly high or low levels of individual transcripts. Such differences in gene expression

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undoubtedly contribute to the observed heterogeneity in biological properties of cancers derived from the same organ.

EXAMPLE 6

This example demonstrates the identities of some of the transcripts which were found to be differentially expressed in tumor and normal tissues. What are the identities of the differentially expressed genes? Of the 548 differentially expressed transcripts, 337 were tentatively identified through database comparisons. When tested, the great majority (93%) of these identifications proved to be legitimate (13), as expected from previous SAGE analyses. Although a large number of differentially expressed genes were identified, some simple patterns did emerge. For example, genes that were expressed at higher levels in normal colon epithelium than in CR tumors were often differentiation-related. These genes included liver fatty acid binding protein, cytokeratin 20, carbonic anhydrase, guanylin and uroguanylin, which are known to be important for the normal physiology or architecture of the colon epithelium (Table 2). On the other hand, genes that were increased in CR cancers were often related to the robust growth characteristics that these cells exhibit. For example, gene products associated with protein synthesis, including 48 ribosomal proteins, five elongation factors, and five genes involved in glycolysis were observed to be elevated in both CR and pancreatic cancers compared to normal colon cells. Although the majority of the transcripts could not have been predicted to be differentially expressed in cancers, several have previously been shown to be dysregulated in neoplastic cells. The latter included IGFII, B23 nucleophosmin, the Pi form of glutathione S-transferase, and several ribosomal proteins which were all increased in cancer cells as previously reported. Likewise, Dra and gelsolin were both decreased in cancer as previously reported. Surprisingly, two widely studied oncogenes, c-fos and c-erbb3, were expressed at much higher levels in normal colon epithelium than CR cancers, in contrast to their up-regulation in transformed cells.

In summary, these data provide basic information necessary for understanding the gene expression differences that underlie cancer phenotypes. They additionally provide a necessary framework for interpreting the significance of individual differentially expressed genes. Although this study demonstrated that a large number of such differences exist (approximately 500 at the depth of analysis employed), it was equally remarkable that the fraction of transcripts exhibiting significant differences was relatively small, representing 1.5 % of the transcripts detected in any given cell type (26). The fact that many, but not all, of the differences were preserved during in vitro culture demonstrates the utility of cultured lines for examination of some aspects of gene expression, but also provides a note of caution in relying on such lines to perfectly mimic tumors in their natural environment. Finally, the finding that hundreds of specific genes are expressed at different levels in CR cancers, and that some of these are also expressed differentially in pancreatic cancers, provides a wealth of new reagents for future biologic and diagnostic experimentation.

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- 3. To minimize individual variation, approximately equal numbers of tags (30,000) were derived from two different patients for each tissue. For primary tumors (two CR carcinomas and two pancreatic adenocarcinomas), RNA was isolated from portions of tumors judged to contain 60%-90% tumor cells by histopathology. The cells grown in vitro were derived from CR (SW837, Caco2) and pancreatic (ASPC-1, PL45) cancer cell lines. CR epithelial cells were isolated from sections of normal colon mucosa from two patients using EDTA as previously described [S. Nakamura, I. Kino, S. Baba, Gut 34, 1240 (1993)]. Histopathology confirmed that the isolated cells were greater than 90% epithelial. Isolation of Poly-A RNA and SAGE was performed as previously described (2). SAGE data was analyzed by means of SAGE software and GenBank Release 95 as previously described (2).
- 4. A total of 69,393 different SAGE tags were identified among the 303,706 tags analyzed. A small fraction of these different tags were likely due to sequencing errors. SAGE analysis of yeast (2), wherein the entire genomic sequence is known, demonstrated a sequencing error rate of ~ 0.7%, translating to a SAGE tag error rate of 6.8% (1 0.993¹⁰). Because these sequencing mistakes are essentially random, they do not substantially affect the analysis although they could artificially inflate the number of unique genes identified. Therefore, to be conservative, we reduced our estimate of unique genes identified by this maximum tag error rate (e.g., 6.8% of 303,706 total tags). The number of different tags derived from the same gene due to alternative splicing was assumed to be negligible.

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- 5. Abundances can be simply determined by dividing the observed number of tags for a given transcript by the total number of tags obtained. An estimate of approximately 300,000 transcripts per cell was used to convert the abundances to copies per cell [N. D. Hastie, J. O. Bishop, *Cell* 9, 761 (1976)].
- J. O. Bishop, J. G. Morton, M. Rosbash, M. Richardson, *Nature* 250, 199 (1974); B. Lewin, Gene Expression Vol 2 (John Wiley and sons, New York 1980).
- 7. Computer simulations indicated that analysis of 300,000 tags would yield a 92 % chance of detecting a tag for a transcript whose expression was at least three copies per cell on average among the tissues examined and assuming 300,000 transcripts per cell.
- 8. To minimize the number of assumptions and to account for the large number of comparisons being made, Monte Carlo analysis was used for determining statistical significance. The null hypothesis was that the level, kind, and distribution of transcripts were the same for cancer and normal cells. For each transcript, 100,000 simulations were performed to determine the relative likelihood due to chance alone ("p-chance") of obtaining a difference in expression equal to or greater than the observed difference, given the null hypothesis. This likelihood was converted to an absolute probability value by simulating 40 experiments in which a representative number of transcripts (27,993 transcripts in each experiment) was identified and compared. The distribution of transcripts used for these simulations was derived from the average level of expression observed in the original samples. The distribution of the p-chance scores obtained in the 40 simulated experiments (false positives) was then compared to those obtained experimentally. Based on this comparison, a maximum value of 0.0005 was chosen for p-chance. This yielded a false positive rate that was no higher than 0.01 for the least significant p-chance value below the cutoff.
- Two hundred simulations assuming an abundance of 0.0001 in one sample and 0.0006 in a second sample revealed a significant difference (P < 0.01, [8]) 95% of the time.

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- 10. It is not possible to obtain pancreatic ductal epithelium, from which pancreatic carcinomas arise, in sufficient quantities to perform SAGE. It is therefore not possible to determine whether these transcripts were derived from genes that were highly expressed only in pancreatic cancers or were also expressed in pancreatic duct cells.
- 11. Total RNA isolation and Northern blot analysis was performed as described [W. S. el-Deiry, et al., Cell 75, 817 (1993)].
- 12. A. H. Owens, D. S. Coffey, S. B. Baylin, Eds., Tumor Cell Heterogeneity: Origins and Implications (Academic Press, New York, 1982).
- 13. Northern blot analyses were done on 45 of the 337 differentially expressed transcripts with tentative database matches. In three cases, the pattern of expression was not differentially expressed as predicted by SAGE and, for the purposes of this calculation, were presumed to represent incorrect database matches.
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 H. Kraus, W. Issing, T. Miki, N. C. Popescu, S. A. Aaronson, Proc Natl Acad Sci USA 86, 9193 (1989).
- 26. In the case of normal and neoplastic colon cancer tissue, 548 differentially transcripts were identified among the 36,125 unique transcripts.
 - 27. All references cited are hereby incorporated by reference herein.
- 28. Sequences tags in Tables 2-4 are consecutively numbered to form SEQ ID NOS: 1-732.

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CLAIMS

1. A method of diagnosing colon cancer in a sample suspected of being neoplastic, comprising the steps of:

comparing the level of at least one transcript in a first sample of a tissue to a second sample, wherein the first sample is of a colonic tissue suspected of being neoplastic and the second sample is of a normal human colonic tissue, and wherein the transcript is identified by a tag selected from the group consisting of those shown in Table 3;

identifying the first sample as neoplastic when the level of the at least one transcript is found to belower in the first sample than in the second sample.

2. A method of diagnosing colon cancer in a sample suspected of being neoplastic, comprising the steps of:

comparing the level of at least one transcript in a first sample of a tissue to a second sample, wherein the first sample is of a colonic tissue suspected of being neoplastic and the second sample is of a normal human colonic tissue, and wherein the transcript is identified by a tag selected from the group consisting of those shown in Table 2;

identifying the first sample as neoplastic when the level of the at least one transcript is found to be higher in the first sample than in the second sample.

- The method of claim 1 wherein a comparison of at least two of said transcripts is performed.
- 4. The method of claim 2 wherein a comparison of at least two of said transcripts is performed.

- 5. The method of claim I wherein a comparison of at least five of said transcripts is performed.
- 6. The method of claim 2 wherein a comparison of at least five of said transcripts is performed.
- The method of claim 1 wherein a comparison of at least ten of said transcripts is performed.
 - 8. The method of claim 2 wherein a comparison of at least ten of said transcripts is performed.
 - 9. The method of claim 1 wherein a comparison of at least twenty of said transcripts is performed.
 - 10. The method of claim 2 wherein a comparison of at least twenty of said transcripts is performed.
 - 11. The method of claim 1 wherein a comparison of at least thirty of said transcripts is performed.
- 15 12. The method of claim 2 wherein a comparison of at least thirty of said transcripts is performed.
 - An isolated and purified human nucleic acid molecule which comprises a SAGE tag selected from SEQ ID NO:1-732.
 - 14. The nucleic acid molecule of claim 13 which is a cDNA molecule.

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- 15. The nucleic acid molecule of claim 13 wherein the SAGE tag is located at the 3' end of the molecule, adjacent to the 3'-most NlaIII restriction enzyme site.
- 16. An isolated nucleotide probe comprising at least 10 nucleotides of a human nucleic acid molecule, wherein the human nucleic acid molecule comprises a SAGE tag selected from SEQ ID NO: 1-732.
- 17. The probe of claim 16 which comprises the selected SAGE tag.
- 18. A diagnostic reagent for evaluating neoplasia of a colorectal tissue, comprising at least 2 probes according to claim 16.
- 19. The diagnostic reagent of claim 18 which comprises at least 5 probes according to claim 16.
 - 20. The diagnostic reagent of claim 18 which comprises at least 10 probes according to claim 16.
 - 21. The diagnostic reagent of claim 18 which comprises at least 20 probes according to claim 16.
 - 22. The diagnostic reagent of claim 18 which comprises at least 30 probes according to claim 16.
 - 23. A diagnostic reagent for evaluating neoplasia of a colorectal tissue, comprising at least 2 probes according to claim 17.
 - 24. A method of diagnosing pancreatic cancer in a sample suspected of being neoplastic, comprising the steps of:

comparing the level of at least one transcript in a first sample of a tissue to a second sample, wherein the first sample is of a pancreatic tissue suspected of being neoplastic and the second sample is of a normal human colon tissue, wherein said transcript is identified by a tag selected from the group consisting of those shown Table 4;

identifying the first sample as neoplastic when the level of the at least one transcript is found to be higher in the first sample than in the second sample.

25. A method of diagnosing cancer in a sample suspected of being neoplastic, comprising the steps of:

comparing the level of at least one transcript in a first sample of a tissue to a second sample, wherein the first sample is of a tissue suspected of being neoplastic and the second sample is of a normal human tissue of the same tissue type, wherein said transcript is identified by a tag selected from the group consisting of those shown Table 5;

identifying the first sample as neoplastic when the level of the at least one transcript is found to be higher in the first sample than in the second sample.

26. A method to aid in the determination of a prognosis for a colon cancer patient, comprising the steps of:

comparing the level of at least one transcript in a first sample of a tissue to a second sample, wherein the first sample is of a neoplastic colonic tissue and the second sample is of a normal human colonic tissue, and wherein the transcript is identified by a tag selected from the group consisting of those shown in Table 3;

determining a poorer prognosis if the level of the at least one transcript is found to be lower in the first sample than in the second sample.

27. A method to aid in determining a prognosis for a patient with colon cancer, comprising the steps of:

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comparing the level of at least one transcript in a first tissue sample to a second sample, wherein the first sample is of a colonic cancer tissue and the second sample is of a normal human colonic tissue, and wherein the transcript is identified by a tag selected from the group consisting of those shown in Table 2;

determining a poorer prognosis if the level of the at least one transcript is found to be higher in the first sample than in the second sample.

28. A method of diagnosing colon cancer in a sample suspected of being neoplastic, comprising the steps of:

comparing the level of expression of at least one protein in a first sample of a tissue to a second sample, wherein the first sample is of a colonic tissue suspected of being neoplastic and the second sample is of a normal human colonic tissue, and wherein the protein is encoded by a transcript identified by a tag selected from the group consisting of those shown in Table 3;

identifying the first sample as neoplastic when the level of expression of the protein is found to be lower in the first sample than in the second sample.

29. A method of diagnosing colon cancer in a sample suspected of being neoplastic, comprising the steps of:

comparing the level of expression of at least one protein in a first sample of a tissue to a second sample, wherein the first sample is of a colonic tissue suspected of being neoplastic and the second sample is of a normal human colonic tissue, and wherein the protein is encoded by a transcript identified by a tag selected from the group consisting of those shown in Table 2;

identifying the first sample as neoplastic when expression of the protein is found to be higher in the first sample than in the second sample.

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30. A method to aid in determining a prognosis of a patient having pancreatic cancer, comprising the steps of:

comparing the level of at least one transcript in a first sample of a tissue to a second sample, wherein the first sample is of a neoplastic pancreatic tissue and the second sample is of a normal human colon tissue, wherein said transcript is identified by a tag selected from the group consisting of those shown Table 4;

determining a poorer prognosis if the level of the at least one transcript is found to be higher in the first sample than in the second sample.

31. A method to aid in providing a prognosis for a cancer patient, comprising the steps of:

comparing the level of at least one transcript in a first sample of a tissue to a second sample, wherein the first sample is of a neoplastic tissue and the second sample is of a normal human tissue of the same tissue type, wherein said transcript is identified by a tag selected from the group consisting of those shown Table 5;

determining a poorer prognosis if the level of the at least one transcript is found to be higher in the first sample than in the second sample.

32. A method of diagnosing pancreatic cancer in a sample suspected of being neoplastic, comprising the steps of:

comparing the level of expression of at least one protein encoded by a transcript in a first sample of a tissue to a second sample, wherein the first sample is of a pancreatic tissue suspected of being neoplastic and the second sample is of a normal human colon tissue, wherein said protein is encoded by a transcript identified by a tag selected from the group consisting of those shown Table 4;

identifying the first sample as neoplastic when expression of the protein is found to be higher in the first sample than in the second sample.

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33. A method of diagnosing cancer in a sample suspected of being neoplastic, comprising the steps of:

comparing the level of expression of at least one protein in a first sample of a tissue to a second sample, wherein the first sample is of a tissue suspected of being neoplastic and the second sample is of a normal human tissue, wherein said protein is encoded by a transcript identified by a tag selected from the group consisting of those shown Table 5;

identifying the first sample as neoplastic when expression of the protein is found to be higher in the first sample than in the second sample.

34. A method to aid in the determination of a prognosis for a colon cancer patient, comprising the steps of:

comparing the level of expression of at least one protein in a first sample of a tissue to a second sample, wherein the first sample is of a neoplastic colonic tissue and the second sample is of a normal human colonic tissue, and wherein the protein is encoded by a transcript identified by a tag selected from the group consisting of those shown in Table 3;

determining a poorer prognosis if the level of expression is found to be lower in the first sample than in the second sample.

35. A method to aid in determining a prognosis for a patient with colon cancer, comprising the steps of:

comparing the level of expression of at least one protein in a first tissue sample to a second sample, wherein the first sample is of a colonic cancer tissue and the second sample is of a normal human colonic tissue, and wherein the protein is encoded by a transcript identified by a tag selected from the group consisting of those shown in Table 2;

determining a poorer prognosis if the level of expression is found to be higher in the first sample than in the second sample.

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36. A method to aid in determining a prognosis of a patient having pancreatic cancer, comprising the steps of:

comparing the level of expression of at least one protein in a first sample of a tissue to a second sample, wherein the first sample is of a neoplastic pancreatic tissue and the second sample is of a normal human colon tissue, wherein said protein is encoded by a transcript identified by a tag selected from the group consisting of those shown Table 4;

determining a poorer prognosis if the level of expression is found to be higher in the first sample than in the second sample.

37. A method to aid in providing a prognosis for a cancer patient, comprising the steps of:

comparing the level of expression of at least one protein in a first sample of a tissue to a second sample, wherein the first sample is of a neoplastic tissue and the second sample is of a normal human tissue of the same tissue type, wherein said protein is encoded by a transcript identified by a tag selected from the group consisting of those shown Table 5;

determining a poorer prognosis if the level of expression is found to be higher in the first sample than in the second sample.

38. A method of treating a cancer cell, comprising the step of:

administering to a cancer cell an antibody which specifically binds to a protein encoded by a transcript identified by a tag selected from the group consisting of those shown in Tables 2, 4, and 5, wherein the antibody is linked to a cytotoxic agent.

39. An antibody linked to a cytotoxic agent, wherein the antibody specifically binds to a protein encoded by a transcript identified by a tag selected from the group consisting of those shown in Tables 2, 4, and 5.

40. A method of detecting colon cancer in a patient, comprising the steps of:

comparing the level of at least one protein in a first body sample to a second body sample, wherein the first sample is a body sample of the patient and the second sample is of a normal human, wherein the protein is encoded by a transcript identified by a tag selected from the group consisting of those shown in Table 2, wherein the first and second body sample is a sample selected from the group consisting of blood, urine, feces, sputum, and serum;

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identifying neoplasia when the level of the at least one protein is found to be higher in the first sample than in the second sample.

A method of detecting pancreatic cancer in a patient, comprising the 41. steps of:

comparing the level of at least one protein encoded by a transcript in a first sample of a tissue to a second sample, wherein the first sample is of the patient and the second sample is of a normal human, wherein said protein is encoded by a transcript identified by a tag selected from the group consisting of those shown Table 4, wherein the first and second sample is a sample selected from the group consisting of blood, urine, feces, sputum, and serum;

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identifying neoplasia when the level of the at least one protein is found to be higher in the first sample than in the second sample.

A method of detecting cancer in a patient, comprising the steps of: comparing the level of at least one protein in a first sample to 25 a second sample, wherein the first sample is of patient and the second sample is of a normal human, wherein said protein is encoded by a transcript identified by a tag selected from the group consisting of those shown Table 5, wherein the first and second body sample is a sample selected from the group consisting

of blood, urine, feces, sputum, and serum;

identifying neoplasia when the level of the at least one protein is found to be higher in the first sample than in the second sample.

43. A method to aid in determining a prognosis for a patient with colon cancer, comprising the steps of:

comparing the level of at least one protein in a first sample to a second sample, wherein the first sample is of a colonic cancer patient and the second sample is of a normal human, wherein the protein is encoded by a transcript identified by a tag selected from the group consisting of those shown in Table 2, wherein the first and second sample is a sample selected from the group consisting of blood, urine, feces, sputum, and serum;

determining a poorer prognosis if the level of the at least one protein is found to be higher in the first sample than in the second sample.

44. A method to aid in determining a prognosis of a patient having pancreatic cancer, comprising the steps of:

comparing the level of at least one protein in a first sample to a second sample, wherein the first sample is of a pancreatic cancer patient and the second sample is of a normal human, wherein said protein is encoded by a transcript identified by a tag selected from the group consisting of those shown Table 4, wherein said first and second sample is a sample selected from the group consisting of blood, urine, feces, sputum, and serum;

determining a poorer prognosis if the level of the at least one protein is found to be higher in the first sample than in the second sample.

45. A method to aid in providing a prognosis for a cancer patient, comprising the steps of:

comparing the level of expression of at least one protein in a first sample to a second sample, wherein the first sample is of a cancer patient and the second sample is of a normal human, wherein said protein is encoded by a transcript identified by a tag selected from the group consisting of those

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shown Table 5, wherein the first and second sample is a sample selected from the group consisting of blood, urine, feces, sputum, and serum;

determining a poorer prognosis if the level of the at least one protein is found to be higher in the first sample than in the second sample.

46. A method of detecting colon cancer in a patient, comprising the steps of:

comparing the level of at least one transcript in a first body sample to a second body sample, wherein the first sample is a body sample of the patient and the second sample is of a normal human, wherein the transcript is identified by a tag selected from the group consisting of those shown in Table 2, wherein the first and second body sample is a sample selected from the group consisting of blood, urine, feces, sputum, and serum;

identifying neoplasia when the level of the at least one transcript is found to be higher in the first sample than in the second sample.

47. A method of detecting pancreatic cancer in a patient, comprising the steps of:

comparing the level of at least one transcript in a first sample of a tissue to a second sample, wherein the first sample is of the patient and the second sample is of a normal human, wherein said transcript is identified by a tag selected from the group consisting of those shown Table 4, wherein the first and second sample is a sample selected from the group consisting of blood, urine, feces, sputum, and serum;

identifying neoplasia when the level of the at least one transcript is found to be higher in the first sample than in the second sample.

48. A method of detecting cancer in a patient, comprising the steps of:

comparing the level of at least one transcript in a first sample to
a second sample, wherein the first sample is of patient and the second sample
is of a normal human, wherein said transcript is identified by a tag selected

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from the group consisting of those shown Table 5, wherein the first and second body sample is a sample selected from the group consisting of blood, urine, feces, sputum, and serum;

identifying neoplasia when the level of the at least one transcript is found to be higher in the first sample than in the second sample.

49. A method to aid in determining a prognosis for a patient with colon cancer, comprising the steps of:

comparing the level of at least one transcript in a first sample to a second sample, wherein the first sample is of a colonic cancer patient and the second sample is of a normal human, wherein the transcript is identified by a tag selected from the group consisting of those shown in Table 2, wherein the first and second sample is a sample selected from the group consisting of blood, urine, feces, sputum, and serum;

determining a poorer prognosis if the level of the at least one transcript is found to be higher in the first sample than in the second sample.

50. A method to aid in determining a prognosis of a patient having pancreatic cancer, comprising the steps of:

comparing the level of at least one transcript in a first sample to a second sample, wherein the first sample is of a pancreatic cancer patient and the second sample is of a normal human, wherein said transcript is identified by a tag selected from the group consisting of those shown Table 4, wherein said first and second sample is a sample selected from the group consisting of blood, urine, feces, sputum, and serum;

determining a poorer prognosis if the level of the at least one transcript is found to be higher in the first sample than in the second sample.

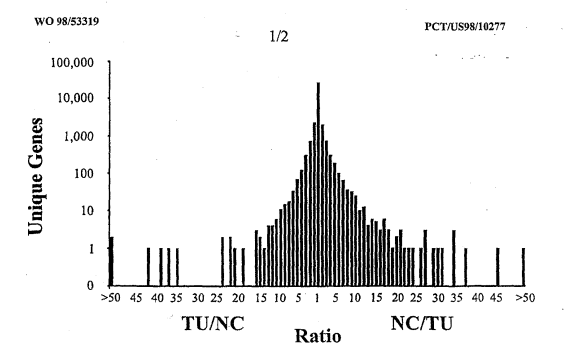
51. A method to aid in providing a prognosis for a cancer patient, comprising the steps of:

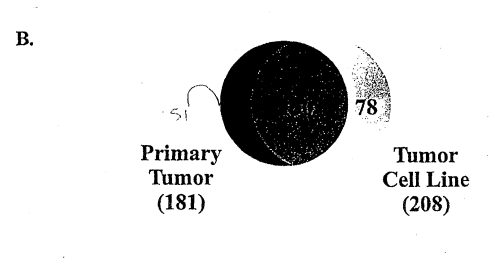
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comparing the level of expression of at least one transcript in a first sample to a second sample, wherein the first sample is of a cancer patient and the second sample is of a normal human, wherein said transcript is identified by a tag selected from the group consisting of those shown Table 5, wherein the first and second sample is a sample selected from the group consisting of blood, urine, feces, sputum, and serum;

determining a poorer prognosis if the level of the at least one transcript is found to be higher in the first sample than in the second sample.

52. A method for screening for candidate agents that modulate the expression of a polynuleotide selected from the group consisting of the polynucleotides in SEQ ID NOS:1-732 or their respective complements, comprising contacting a test agent with a colon or pancreatic cell and monitoring expression of the polynucleotide, wherein the test agent which modifies the expression of the polynucleotide is a candidate agent.





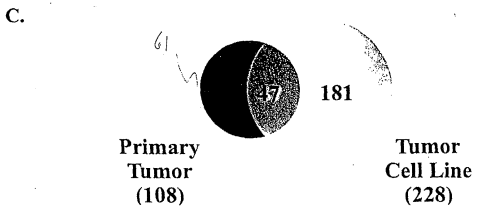


FIG. 2

A.

	1	2	SAGE Data		
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H204104				11	102
H259108	•			1	37
H1000193	041) w (to constitution of the con	56	12
H998030	(1) ()	55	7

B.

			-	ancre Tum						mal ion	SAGE Data		
	1	2	3	4	5	6	7	8	1	2	Pancreatic Tumors	Normal Colon	
				H						Н	Iumors	Colon	
H294155	•	teggi	-	e ingl				The second			47	0	
H560056				-					į		32	0	

C.

	CR Tumors		Pancreatic Tumors			Normal Colon			SAGE Data			
	1	2	3		2		1	2	3	CR Tumors	Pancreatic Tumors	Normal Colon
H802810										27	0	1
H85882		-		•	•) .		* .	10	26	0
H618841				-		-	,	,		8	62	0